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The Flavor and Fragrance High Production Volume Consortia

The Terpene Consortium

Revised Robust Summaries for Monoterpene Hydrocarbons

<i>d</i> -Limonene	CAS No. 5989-27-5
<i>dl</i> -Limonene	CAS No. 138-86-3
Terpinolene	CAS No. 586-62-9
Myrcene	CAS No. 123-35-3
Dihydromyrcene	CAS No. 2436-90-0
Hydrocarbons, terpene processing by-products	CAS No. 68956-56-9
Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck)	CAS No. 8008-57-9
Terpenes & terpenoids, sweet orange oil	CAS No. 68647-72-3
Terpenes & terpenoids, turpentine oil, limonene fraction	CAS No. 65996-99-8
Terpenes & terpenoids, limonene fraction	CAS No. 65996-98-7
Terpenes & terpenoids, turpentine oil, limonene fraction, distillation residue	CAS No. 68334-40-7
Terpenes & terpenoids, turpentine-oil residue	CAS No. 68938-00-1

FFHPVC Terpene Consortium Registration Number

Submitted to the EPA under the HPV Challenge Program by:

The Flavor and Fragrance High Production Volume Chemical Consortia

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Summary of Key Hazard Data for Monoterpene Hydrocarbons

Endpoint	Substance/Surrogate ¹	Value/Range ²	Reference
Physical Properties			
Partition Coefficient	Terpinolene	5.3 (OECD 117)	Givaudan Roure Inc. ,1996a
Partition Coefficient	Terpenes & terpenoids, sweet orange oil	5.3 (OECD 117)	Givaudan Roure Inc. ,1996a
Environmental Fate			
Biodegradation	Terpinolene	28d/62.1%/(OECD 301B)	Birch R., 1996
Biodegradation	Terpinolene	28d/80%/(OECD 302C)	Rudio J., 1998
Biodegradation	Terpinolene	28d/51%/(OECD 301F)	Rudio J. ,1997
Ecotoxicity			
Fish	<i>d</i> -Limonene	96-hr/LC50=0.702 mg/L	Broderius et al., 1990
Fish	<i>d</i> -Limonene	96-hr/LC50=0.720 mg/L	Broderius et al., 1990
Fish	Terpinolene	96-hr/LC50=1.210 mg/L	Broderius et al., 1990
Aquatic Invertebrates	<i>d</i> -Limonene	48-hour LC50 = 0.577 mg/L	Broderius et al., 1990
Aquatic Invertebrates	<i>d</i> -Limonene	48-hour LC50 = 0.924 mg/L	Broderius et al., 1990
Aquatic Invertebrates	Terpinolene	48-hour LC50 = 2.55 mg/L	Broderius et al., 1990
Aquatic Plants	<i>d</i> -Limonene	No significant inhibition	Broderius et al., 1990
Aquatic Plants	Terpinolene	No significant inhibition	Broderius et al., 1990
Human Health			
Repeat Dose (route)	<i>d</i> -Limonene (oral-gavage)	103 wks LOAEL: 75 mg/kg bw/d (male); 600 mg/kg bw/d (female) 103 wks NOAEL=300 mg/kg bw/d (female rat); undetermined	NTP, 1990

¹ Surrogate is a structurally related substance include a metabolic product or precursor of the named substance

² Experimental value or values for a substance or group of substances in the chemical category

		(male rat)	
Repeat Dose (route)	<i>d</i> -Limonene (oral-gavage)	103 wks LOAEL: Undetermined for males; 1000 mg/kg bw/d for female 103 wks NOAEL=500 mg/kg bw/d	NTP, 1990
Repeat Dose (route)	<i>d</i> -Limonene (oral-gavage)	16 d LOEL=3300 mg/kg bw/d 16 d NOEL=1650 mg/kg	NTP, 1990
Repeat Dose (route)	<i>d</i> -Limonene (oral-gavage)	13 wk LOEL=600 mg/kg bw/d 13 wk NOEL=300 mg/kg	NTP, 1990
Repeat Dose (route)	<i>d</i> -Limonene (oral-gavage)	13 wk LOEL=500 mg/kg bw/d 13 wk NOEL=1000 mg/kg	NTP, 1990
Repeat Dose (route)	Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck) (oral-gavage)	28 d LOEL = 240 mg/kg bw/d 28 d NOEL = Not determined	Serota, 1990
Repeat Dose (route)	<i>beta</i> -Myrcene (oral-gavage)	13-week LOEL = 500 mg/kg bw/d 13-wk NOEL = Not determined	NTP draft, 2003
Repeat Dose (route)	<i>beta</i> -Myrcene (oral-gavage)	13-week LOEL= 500 mg/kg bw/d (female) 13-wk NOEL = 250 mg/kg bw/d 13-week LOEL = 1000 mg/kg bw/d (male) 13-wk NOEL= 500 mg/kg bw/d (male)	NTP draft, 2003
Reproductive	<i>beta</i> -Myrcene (oral-gavage)	86-112 d NOEL= 300 mg/kg bw/d LOEL=500 mg/kg bw/d	Paumgarten et al., 1998
e) Developmental(rout	<i>beta</i> -Myrcene (oral-gavage)	128 d NOEL (rat): 250 mg/kg bw/d 128 d LOEL(rat): 500 mg/kg bw/d (oral- gavage)	Delgado et al., 1993b
in vitro Genotoxicity³	<i>d</i> -Limonene; <i>beta</i> -Myrcene; Sweet Orange Oil	-(AMS); - /+(MLA); -, (ABS); - (SCE);	Heck et al., 1989; Florin et al., 1980; Muller, 1993; Haworth et al., 1983; Anderson et

³ (-), no significant evidence; (+/-), equivocal evidence; (+), positive evidence of genotoxicity

			al., 1990; Myhr et al., 1990; Kauderer et al, 1991; Roscheisen et al, 1991; Crebelli et al., 1990; Kuroda et al., 1989.
<i>in vivo Genotoxicity</i>	d-Limonene beta-Myrcene	(-) mouse embryo (-/+) ABS (-) MN	Fahrig, 1984; Zamith et al., 1993. NTP, 2003.

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The evaluation of the quality of the following data uses a systematic approach described by Klimisch [Klimisch *et al.*, 1996]. Based on criteria relating to international testing standards for categorizing data reliability, four reliability categories have been established. The following categories are:

- Reliability code 1. Reliable without restrictions
- Reliability code 2. Reliable with restrictions
- Reliability code 3. Not reliable
- Reliability code 4. Not assignable

1 Chemical and Physical Properties

1.1 Melting Point

Substance Name	d-Limonene
CAS No.	5989-27-5
Method/guideline	Measured
Melting Point	-74.35 °C
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, Florida.

Substance Name	dl-Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Measured

Melting Point	-97 °C
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, Florida.

1.2 Boiling Point

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	176 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Fragrance Materials Association (FMA) Boiling point.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	178.6 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
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CAS No.	5989-27-5
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	175.5-176 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Merck Index (1996) 12th edition, Susan Budavari, editor, Merck & Co. Inc. Whitehouse Station, NJ.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	175.5-176.5 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Merck Index (1996) 12th edition, Susan Budavari, editor, Merck & Co. Inc. Whitehouse Station, NJ.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	178.6 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, FL.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	178 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Fragrance Materials Association (FMA) Boiling point.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	185 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Fragrance Materials Association (FMA) Boiling point.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	185.8 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	172 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Fragrance Materials Association (FMA) Boiling point.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	167.7 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	44 °C
Pressure	10 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.

References

Merck Index (1996) 12th edition, Susan Budavari, editor, Merck & Co. Inc. Whitehouse Station, NJ.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	165-168 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	CRC Handbook of Chemistry and Physics (1986) 67th edition, Robert C. Weast, editor, The Chemical Rubber Co Press, Inc. Boca Raton, FL.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	158 °C
Pressure	760 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Fragrance Materials Association (FMA) Boiling point.

Substance Name	Terpenes & terpenoids, sweet orange oil
CAS No.	68647-72-3
Method/guideline	Measured
GLP	Ambiguous
Boiling Point	176 °C
Pressure	1013 mm Hg
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Basic data given: comparable to guidelines/standards.

References Givaudan-Roure (1991) Unpublished report to RIFM.

1.3 Vapor Pressure

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Measured
Vapor Pressure	1.43 mmHg (0.19 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Only secondary literature (review, tables, books, etc.).
References	IPCS (1998) Concise International Chemical Assessment Document No. 5 Limonene. World Health Organization, 5, Geneva.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Calculated
Vapor Pressure	1.2 mm Hg (0.16 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	Fragrance Materials Association (FMA) Report values for vapor pressure.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Calculated/Mean of Antoine & Grain
Vapor Pressure	1.59 mmHg (0.19 kPa)
Temperature	20 °C

Data Qualities Reliabilities Reliability code 4. Not assignable
Remarks for Data Reliability Code 4. Calculated.
References MPBPVPWIN EPI suite (2000) US Environmental Protection Agency.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Calculated/Mean of Antoine & Grain
Vapor Pressure	1.63 mm Hg (0.22 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	MPBPVPWIN EPI suite (2000) US Environmental Protection Agency.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Calculated
Vapor Pressure	0.8 mm Hg (0.1 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	Fragrance Materials Association (FMA) Report values for vapor pressure.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Calculated/Mean of Antoine & Grain
Vapor Pressure	1.44 mm Hg (0.19 kPa)
Temperature	25 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.

References MPBPVPWIN EPI suite (2000) US Environmental Protection Agency.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Calculated
Vapor Pressure	0.5 mm Hg (0.07 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	Fragrance Materials Association (FMA) Report values for vapor pressure.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Measured
Vapor Pressure	2.01 mm Hg
Temperature	25 oC
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).
References	Perry, R.H.; Green, D. 1984. Perry's Chemical Handbook. Physical and Chemical Data, New York, NY: McGraw-Hill, 6th ed.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Calculated
Vapor Pressure	1.5 mm Hg (0.2 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	Fragrance Materials Association (FMA) Report values for vapor pressure.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Calculated/Mean of Antoine & Grain
Vapor Pressure	2.4 mm Hg (0.32 kPa)
Temperature	25 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	MPBPVPWIN EPI suite (2000) US Environmental Protection Agency.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	Calculated/Mean of Antoine & Grain
Vapor Pressure	2.57 mm Hg (0.34 kPa)
Temperature	25 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	MPBPVPWIN EPI suite (2000) US Environmental Protection Agency.

Substance Name	Orange peel oil, sweet (Citrus sinensis (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	Calculated
Vapor Pressure	0.9 mm Hg (0.12 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	Fragrance Materials Association (FMA) Report values for vapor pressure.

Substance Name	Terpenes & terpenoids, sweet orange oil
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CAS No.	68647-72-3
Method/guideline	Calculated
Vapor Pressure	0.9 mm Hg (0.12 kPa)
Temperature	20 °C
Data Qualities Reliabilities	Reliability code 4. Not assignable
Remarks for Data Reliability	Code 4. Calculated.
References	Fragrance Materials Association (FMA) Report values for vapor pressure.

1.4 n-Octanol/Water Partition Coefficient

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Measured
GLP	Ambiguous
Log Pow	4.23
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Only secondary literature (review, tables, books, etc.).
References	IPCS (1998) Concise International Chemical Assessment Document No. 5 Limonene. World Health Organization, 5, Geneva.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	OECD Guideline No. 117
GLP	Yes
Year	1996
Log Pow	5.3
Temperature	30 °C
Remarks for Test Conditions	Reverse phase HPLC
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Guideline study.

References Givaudan Roure Inc. (1996a) Partition coefficient n-octanol/water of terpinolene. Private communication to RIFM.

Substance Name	Terpenes & terpenoids, sweet orange oil
CAS No.	68647-72-3
Method/guideline	OECD Guideline No. 117
GLP	Yes
Year	1996
Log Pow	5.3
Temperature	45 °C
Remarks for Test Conditions	Reverse phase HPLC
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	Givaudan Roure Inc. (1996b) Partition coefficient n-octanol/water of orange peel. Private communication to RIFM.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Calculated
Log Pow	4.83
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	Syracuse Research Corporation (SRC) Private communication to FMA.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Calculated
Log Pow	4.83
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.

References Syracuse Research Corporation (SRC) Private communication to FMA.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Calculated
Log Pow	4.88
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	Syracuse Research Corporation (SRC) Private communication to FMA.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Calculated
Log Pow	4.88
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	Syracuse Research Corporation (SRC) Private communication to FMA.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	Calculated
Log Pow	4.88
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	Syracuse Research Corporation (SRC) Private communication to FMA.

1.5 Water Solubility

Substance Name	<i>d</i> -Limonene
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CAS No.	5989-27-5
Method/guideline	Experimental
Value (mg/L) at Temperature	13.8 mg/L at 25 °C
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Comparable to guideline study.
References	WSKOWWIN EPI Suite (2000c) US Environmental Protection Agency (Massaldi, HA & King, CJ, 1973).

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Experimental
Value (mg/L) at Temperature	13.8 mg/L at 25 °C
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Comparable to guideline study.
References	WSKOWWIN EPI Suite (2000c) US Environmental Protection Agency (Massaldi, HA & King, CJ, 1973).

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Experimental
Value (mg/L) at Temperature	9.5 mg/L at 23 °C
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Comparable to guideline study.
References	WSKOWWIN EPI Suite (2000b) US Environmental Protection Agency (Li, J. and Perdue, EM, 1995).

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Experimental
Value (mg/L) at Temperature	5.6 mg/L at 25 °C
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Comparable to guideline study.

References

WSKOWWIN EPI Suite (2000a) US Environmental Protection Agency (Chem Inspect Test Inst, 1992).

Substance Name	Terpenes & terpenoids, sweet orange oil
CAS No.	68647-72-3
Method/guideline	Experimental
Value (mg/L) at Temperature	30 mg/L at 20 °C
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Comparable to guideline study with acceptable restrictions.
References	Givaudan-Roure (1991) Unpublished report to RIFM.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Calculated
Value (mg/L) at Temperature	0.74 mg/L at 25 °C
Remarks for Test Conditions	Input parameters: Log Kow, 5.3; Melting Point, -74.35
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	WSKOWWIN EPI Suite (2000d) US Environmental Protection Agency.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Calculated
Value (mg/L) at Temperature	1.87mg/L at 25 °C
Remarks for Test Conditions	Input parameters: Log Kow, 4.88; Melting Point, -29.5
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	WSKOWWIN EPI Suite (2000d) US Environmental Protection Agency.

Substance Name	Myrcene
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CAS No.	123-35-3
Method/guideline	Calculated
Value (mg/L) at Temperature	1.87 mg/L at 25 °C
Remarks for Test Conditions	Input parameters: Log Kow, 4.88; Melting Point, -64.83
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	WSKOWWIN EPI Suite (2000d) US Environmental Protection Agency.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	Calculated
Value (mg/L) at Temperature	1.87 mg/L at 25 °C
Remarks for Test Conditions	Input parameters: Log Kow, 4.88; Melting Point, -66.0
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	WSKOWWIN EPI Suite (2000d) US Environmental Protection Agency.

Substance Name	<i>d</i> -Limonene (dipentene)
CAS No.	138-86-3
Method/guideline	Calculated
Value (mg/L) at Temperature	0.74 mg/L at 25 °C
Remarks for Test Conditions	Input parameters: Log Kow, 5.3; Melting Point, -97
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	WSKOWWIN EPI Suite (2000d) US Environmental Protection Agency.

2 Environmental Fate and Pathways

2.1 Photodegradation

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Calculated
Test Type	AOPWIN
Half-life t_{1/2}	0.884 hours
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	AOPWIN EPI Suite (2000) US Environmental Protection Agency.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Calculated
Test Type	AOPWIN
Half-life t_{1/2}	0.64 hours
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
References	AOPWIN EPI Suite (2000) US Environmental Protection Agency.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Calculated
Test Type	AOPWIN
Half-life t_{1/2}	0.66 hours
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.

References AOPWIN EPI Suite (2000) US Environmental Protection Agency.

2.2 Biodegradation

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	97% purity
Method	Not given
Test Type	Sealed vessel
GLP	Ambiguous
Year	1996
Contact Time	30 days
Innoculum	Unacclimated soil from coniferous forest
Remarks for Test Conditions	Limonene was tested in concentrations of 0.5-3 mg/liter, in sealed reactors. After 24 hours equilibration, inocula were added at 1%. Incubation took place in the dark at 23 degrees Celsius with continuous mixing. Duplicate gas and liquid samples were taken at regular intervals and analyzed for monoterpenes and CO ₂ .
Degradation % After Time	0.044 mg/l h maximum degradation rate
Time required for 10% degradation	Less than 1 day
Results	Limonene was depleted below detectable limits within 1 day in acclimated soil and 8 days in unacclimated soil.
Classification	Readily biodegradable
Conclusion Remarks	The increase in biomass and headspace CO ₂ parallel the depletion of limonene, confirming that the disappearance of limonene was a result of biodegradation.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
Reference	Misra G., Pavlostathis S., Perdue E., Araujo R. (1996) Aerobic biodegradation of selected monoterpenes. Appl Microbiol Biotechnol 45: 831-838.

Substance Name	Terpinolene
CAS No.	586-62-9
Method	OECD Method 301B

Test Type	Sealed vessel carbon dioxide production test
GLP	Yes
Year	1996
Contact Time	28 days
Innoculum	Secondary effluent from an unacclimatized activated sludge plant
Remarks for Test Conditions	Nominal carbon concentrations of the test substances were used based on the calculated percentage carbon and assuming 100% purity of the named compound (1062 micrograms of carbon for terpinolene). The testing procedure followed OECD 301B.
Degradation % After Time	62.1% (95% C.I. 33.3-90.8) after 28 days
Time required for 10% degradation	Less than 3 days
Remarks Results	The air temperature during the 28 day test period was 19-24 degrees Celsius.
10 day window criteria	63.6%
Classification	PASS- readily and ultimately biodegradable
Conclusion Remarks	The test substance, terpinolene, was considered ultimately and readily biodegradable.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Comparable to guideline study.
Reference	Birch R.R. (1996) The ultimate biodegradability of terpinolene in the Sealed Vessel Test. Private Communication to FFHPVC.

Substance Name	Terpinolene
CAS No.	586-62-9
Remarks for Substance	82.2% pure by GC; almost colorless to pale yellow liquid; density 0.875 g/ml at 20 degrees Celsius
Method	OECD Method 302C
Test Type	Inherent Biodegradability: Manometric Respirometry Test
GLP	Yes
Year	1998
Contact Time	31 days
Innoculum	Fresh activated sludge
Remarks for Test Conditions	Followed OECD Method No. 302C. The reference substance used was sodium benzoate. The concentration of test substance used was 30 mg/l. The test temperature was 25 degrees Celsius.

Degradation % After Time	80% after 31 days (also 80% after 28 days)
Time required for 10% degradation	Less than 5 days
Results	The % degradation (mean of 2 identical flasks) after 5 days is 52%; 7 days is 54%; 14 days is 74%; 21 days is 80%; 28 days is 80% and 31 days is 80%.
10 day window criteria	Not given
Classification	Inherently biodegradable
Breakdown products (transient or stable)	Not given
Remarks fields for results	Averages of 2 identical flasks were used to determine the results. Degradation of sodium benzoate was 65% after 7 days and 86% after 14 days; the activity of the inoculum thus verified.
Conclusion Remarks	The test substance, terpinolene, underwent 80% biodegradation after 31 days under the test conditions and is considered inherently biodegradable.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
Reference	Rudio J. (1998) Inherent biodegradability of terpinolene according to OECD Guideline No. 302C. Private communication to FFHPVC.

Substance Name	Terpinolene
CAS No.	586-62-9
Method	OECD Method 301F
Test Type	Ready Biodegradability: Manometric Respirometry Test
GLP	Yes
Year	1997
Contact Time	28 days
Innoculum	Fresh activated sludge
Remarks for Test Conditions	Followed OECD Method No. 301F. The reference substance used was sodium benzoate. The concentration of test substance used was 100 mg/l and the test temperature was 22 degrees Celsius.
Degradation % After Time	51% degradation after 28 days
Time required for 10% degradation	Less than 2 days
Results	The % degradation (mean of 2 identical flasks) after 2 days is 11%; 7 days is 31%; 12 days is 47%; 14 days is 49%; 21 days is 50% and 28 days is 51%.
10 day window criteria	47% at end of 10 day window (days 2 to 12)

Classification	Not readily biodegradable
Breakdown products (transient or stable)	Not given
Remarks fields for results	No toxic effects of terpinolene were observed on the microorganisms.
Conclusion Remarks	Terpinolene, which underwent 51% biodegradation after 28 days, should be regarded as not readily biodegradable under the conditions outlined for this test.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
Reference	Rudio J. (1997) Ready biodegradability of pinene <i>alpha</i> according to OECD Guideline No. 301F. Private communication to FFHPVC.

Substance Name	Terpinolene
CAS No.	586-62-9
Remarks for Substance	97% purity
Method	Not given
Test Type	Sealed vessel
GLP	Ambiguous
Year	1996
Contact Time	30 days
Innoculum	Unacclimated soil from coniferous forest
Remarks for Test Conditions	Terpinolene was tested in concentrations of 0.5-3 mg/liter, in sealed reactors. After 24 hours equilibration, inocula were added at 1%. Incubation took place in the dark at 23 degrees Celsius with continuous mixing. Duplicate gas and liquid samples were taken at regular intervals and analyzed for monoterpenes and CO ₂ .
Degradation % After Time	0.053 mg/l h maximum degradation rate
Time required for 10% degradation	Less than 1 day
Results	Limonene was depleted below detectable limits within 1 day in acclimated soil and 8 days in unacclimated soil.
Classification	Readily biodegradable
Conclusion Remarks	The increase in biomass and headspace CO ₂ parallel the depletion of terpinolene, confirming that the disappearance of terpinolene was a result of biodegradation.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.

Reference

Misra G., Pavlostathis S., Perdue E., Araujo R. (1996) Aerobic biodegradation of selected monoterpenes. Appl Microbiol Biotechnol 45: 831-838.

2.3 Fugacity

Substance	d-Limonene
CAS	5989-27-5
Model Conditions	25 C, 1000 lbs.
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used (title, version, date)	EQC Fugacity Level III
Input parameters	MW (114g/mole), VP(1.43 mm Hg), log Kow (4.23), water solubility (13.8 mg/L), BP (176) estimated MP (-97)
Year	2000
Media	Air-Water-Soil-Sediment Partition Coefficients
Model data and results	Compartment half-lives, hours:Air=0.334; Water=360;Soil=360;Sediment=1440
Estimated Distribution and Media Concentration	Air=0.145% Water=32.1% Soil=64.4% Sediment=3.38%
Conclusion remarks	Substance is predicted to persist in the environment for 192 hours. Persistence data consistent with a measured biodegradation rate in active soil samples of 100% within 1 days.
Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	The data are obtained by a recognized fugacity calculation method. Data are considered reliable with restriction because this method does not allow for biodegradation or metabolism.
References	Mackay D., A.DiGuardo, S.Paterson and C.E.Cowan (1996b) Evaluating the fate of a variety of types of chemicals using the EQC model. Environmental Toxicology and Chemistry, 15(9), 1627-1637.
Substance	Terpinolene
CAS	586-62-9
Model Conditions	25 C, 1000 lbs.
Test Type	Environmental Equilibrium Partitioning Model

Method	Mackay
Model Used (title, version, date)	EQC Fugacity Level III
Input parameters	MW (114g/mole), VP(1.44 mm Hg), log Kow (5.3), water solubility (9.5 mg/L), BP (185) estimated MP
Year	2000
Media	Air-Water-Soil-Sediment Partition Coefficients
Model data and results	Compartment half-lives, hours:Air=0.147; Water=360;Soil=360;Sediment=1440
Estimated Distribution and Media Concentration	Air=0.034% Water=20.3% Soil=58.4% Sediment=21.2%
Conclusion remarks	Substance is predicted to persist in the environment for 292 hours. Persistence data consistent with a measured biodegradation rate in active soil samples of 80% within 31 days.
Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	The data are obtained by a recognized fugacity calculation method. Data are considered reliable with restriction because this method does not allow for biodegradation or metabolism.
References	Mackay D., A.DiGuardo, S.Paterson and C.E.Cowan (1996b) Evaluating the fate of a variety of types of chemicals using the EQC model. Environmental Toxicology and Chemistry, 15(9), 1627-1637.
Substance	Myrcene
CAS	123-35-3
Model Conditions	25 C, 1000 lbs.
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used (title, version, date)	EQC Fugacity Level III
Input parameters	MW (114g/mole), VP(2.01 mm Hg), log Kow (4.88-calc), water solubility (5.6 mg/L), BP (172) estimated MP
Year	2000
Media	Air-Water-Soil-Sediment Partition Coefficients
Model data and results	Compartment half-lives, hours:Air=0.186; Water=360;Soil=360;Sediment=1440
Estimated Distribution and Media Concentration	Air=0.0606% Water=25.9% Soil=62.8% Sediment=11.2%

Conclusion remarks	Substance is predicted to persist in the environment for 233 hours. Persistence data consistent with a measured biodegradation rate in active soil samples of 89% within 25 days (Primary Survey Model).
Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	The data are obtained by a recognized fugacity calculation method. Data are considered reliable with restriction because this method does not allow for biodegradation or metabolism.
References	Mackay D., A.DiGuardo, S.Paterson and C.E.Cowan (1996b) Evaluating the fate of a variety of types of chemicals using the EQC model. Environmental Toxicology and Chemistry, 15(9), 1627-1637.
Substance	Dihydromyrcene

CAS	5989-27-5
Model Conditions	25 C, 1000 lbs.
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used (title, version, date)	EQC Fugacity Level III
Input parameters	MW (114g/mole), VP(2.57 mm Hg-calc), log Kow (4.88-calc), water solubility (1.87 mg/L-calc), BP (167) estimated MP
Year	2000
Media	Air-Water-Soil-Sediment Partition Coefficients
Model data and results	Compartment half-lives, hours:Air=0.483; Water=360;Soil=360;Sediment=1440
Estimated Distribution and Media Concentration	Air=0.226% Water=32.1% Soil=53.8% Sediment=13.9%
Conclusion remarks	Substance is predicted to persist in the environment for 192 hours. Persistence data consistent with a measured biodegradation rate in active soil samples of 100% within 25 days (Primary Survey Model).
Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	The data are obtained by a recognized fugacity calculation method. Data are considered reliable with restriction because this method does not allow for biodegradation or metabolism.
References	Mackay D., A.DiGuardo, S.Paterson and C.E.Cowan (1996b) Evaluating the fate of a variety of types of chemicals using the EQC model. Environmental Toxicology and Chemistry, 15(9), 1627-1637.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Soil-Water Partition Coefficient
Absorption coefficient	731
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Sediment-Water Partition Coefficient
Absorption coefficient	1462
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Suspended Sediment-Water Partition Coefficient
Absorption coefficient	4570
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Air-Water Partition Coefficient
Absorption coefficient	1.05
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Fish-Water Partition Coefficient
Absorption coefficient	1858
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Aerosol-Air Partition Coefficient
Absorption coefficient	22700
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Air
Estimated Distribution and Media Concentration	93.8%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Water
Estimated Distribution and Media Concentration	0.18%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Soil
Estimated Distribution and Media Concentration	5.87%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Sediment
Estimated Distribution and Media Concentration	0.13%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Suspended Sediment
Estimated Distribution and Media Concentration	0.0041%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Fish
Estimated Distribution and Media Concentration	0.00033%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Aerosol
Absorption coefficient	0.000043%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Air-Water Partition Coefficient
Absorption coefficient	0.57
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Sediment-Water Partition Coefficient
Absorption coefficient	1162
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Suspended Sediment-Water Partition Coefficient
Absorption coefficient	3630
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, calculated MP, water solubility
Media	Soil-Water Partition Coefficient
Absorption coefficient	581
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, VP, log Kow, MP, water solubility
Media	Fish-Water Partition Coefficient
Absorption coefficient	1477
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Aerosol-Air Partition Coefficient
Absorption coefficient	60600
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Air
Estimated Distribution and Media Concentration	91.2%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Water
Estimated Distribution and Media Concentration	0.32%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Soil
Estimated Distribution and Media Concentration	8.32%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Sediment
Estimated Distribution and Media Concentration	0.18%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Suspended Sediment
Estimated Distribution and Media Concentration	0.0058%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Fish
Estimated Distribution and Media Concentration	0.00047%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Terpinolene
CAS No.	586-62-9
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Aerosol
Estimated Distribution and Media Concentration	0.000011%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Air-Water Partition Coefficient
Absorption coefficient	2.64
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Soil-Water Partition Coefficient
Absorption coefficient	291
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Sediment-Water Partition Coefficient
Absorption coefficient	582
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Suspended Sediment-Water Partition Coefficient
absorption coefficient	1820
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Fish-Water Partition Coefficient
Absorption coefficient	740
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Aerosol-Air Partition Coefficient
Absorption coefficient	22300
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Air
Estimated Distribution and Media Concentration	98.9%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Water
Estimated Distribution and Media Concentration	0.075%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Soil
Estimated Distribution and Media Concentration	0.098%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Sediment
Estimated Distribution and Media Concentration	0.022%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Suspended Sediment
Estimated Distribution and Media Concentration	0.00068%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Fish
Estimated Distribution and Media Concentration	0.000056%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	Myrcene
CAS No.	123-35-3
Model Conditions	25 °C, 100,000
Test Type	Environmental Equilibrium Partitioning Model
Method	Mackay
Model Used	EQC V 2.11 Level I
Input Parameters	MW, log Kow, water solubility, VP & calculated MP
Media	Aerosol
Estimated Distribution and Media Concentration	0.000044%
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated. The data are obtained by a recognized fugacity calculation method.
References	Trent University (1999) Level 1 Fugacity-based Environmental Equilibrium Partitioning Model Version 2.11. Based on Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

3 Ecotoxicity

3.1 Acute Toxicity to Fish

Substance Name	<i>d</i> -Limonene																																															
CAS No.	5989-27-5																																															
Remarks for Substance	<i>d</i> -Limonene in neat form had purity of 99%, in toxicant saturated water of 85%, in exposure water 67%																																															
Method/guideline	Flow-through. Calculated LC50 and EC50 using the Trimmed Spearman-Karber Method [Hamilton <i>et al.</i> , 1977] and corrected average of the analyzed tank concentrations																																															
Test Type	Experimental																																															
GLP	Ambiguous																																															
Year	1990																																															
Species/Strain/Supplier	Fathead minnows/Fluorostat Corporation																																															
Exposure Period	96 hour																																															
Analytical monitoring	GC Analysis																																															
Remarks for Test Conditions	<p>96 hour LC50 and EC50 tests were performed with fathead minnows using <i>d</i>-limonene from two different suppliers. Test protocol was a continuous flow-through system. Tests were conducted in the electronic diluter using 0.2 L test chambers. A saturator flow rate of 21 mls/min resulted in exposure concentrations of 178 to 1350 ug/L.</p> <p>The ecotoxicity test in fish was conducted with a modified Benoit diluter (Benoit, 1981) which utilized six one-liter tanks; one control and five treatments with a dilution ratio of 0.65 between each treatment. The flow to each tank was 10 ml/min giving 14.4 volume additions per day. One control and five dilutions at 100, 80, 60, 40 and 20 percent of the stock concentration comprised the exposure series. The test solution depth was measured at 4-5 cm. No replicates were used. Tanks were placed in a random sequence in a constant temperature water bath. The water temperature was 24.4 deg C. The pH was 7.6. The dissolved oxygen was 5.8 mg/L. Hardness was 46.2 mg/L CaCO₃ with an alkalinity of 39.5 mg/L CaCO₃.</p>																																															
Endpoint value	<p>LC50 = 702 ug/L 95% C.L. (619-796 ug/L)</p> <p>Mortality</p> <table border="1"> <thead> <tr> <th>Hour</th> <th>Concentration</th> <th>A</th> <th>B</th> <th>C</th> <th>D</th> <th>E</th> </tr> </thead> <tbody> <tr> <td>Initial</td> <td>10</td> <td>10</td> <td>10</td> <td>10</td> <td>10</td> <td>10</td> </tr> <tr> <td>24</td> <td></td> <td></td> <td></td> <td>2</td> <td>9</td> <td>6</td> </tr> <tr> <td>48</td> <td></td> <td></td> <td></td> <td>2</td> <td>9</td> <td>8</td> </tr> <tr> <td>72</td> <td></td> <td></td> <td></td> <td>2</td> <td>9</td> <td>10</td> </tr> <tr> <td>96</td> <td></td> <td></td> <td></td> <td>2</td> <td>9</td> <td>10</td> </tr> </tbody> </table>						Hour	Concentration	A	B	C	D	E	Initial	10	10	10	10	10	10	24				2	9	6	48				2	9	8	72				2	9	10	96				2	9	10
Hour	Concentration	A	B	C	D	E																																										
Initial	10	10	10	10	10	10																																										
24				2	9	6																																										
48				2	9	8																																										
72				2	9	10																																										
96				2	9	10																																										

EC50=702 ug/L 95% C.L(619-796 ug/L)

Mortality

Hour	Concentration	A	B	C	D	E
Initial	10	10	10	10	10	10
24				10	10	10
48				2	10	10
72				2	10	10
96				2	9	10

Measured concentrations as mg/L 2.73 mg/L

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Guideline study.

Reference Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	<i>d</i> -Limonene in neat form had purity of 99%, in toxicant saturated water of 85%, in exposure water 67%
Method/guideline	Flow-through. Calculated LC50 and EC50 using the Trimmed Spearman-Kärber Method [Hamilton <i>et al.</i> , 1977] and corrected average of the analyzed tank concentrations
Test Type	Experimental
GLP	Ambiguous
Year	1990
Species/Strain/Supplier	Fathead minnows/Aldrich Chemical
Exposure Period	96 hour
Analytical monitoring	GC Analysis
Remarks for Test Conditions	<p>96 hour LC50 and EC50 tests were performed with fathead minnows using <i>d</i>-limonene from two different suppliers. Test protocol was a continuous flow-through system. Tests were conducted in the electronic diluter using 0.2 L test chambers. A saturator flow rate of 21 ml/min resulted in exposure concentrations of 251 to 1890 ug/L.</p> <p>The ecotoxicity test in fish was conducted with a modified Benoit diluter (Benoit, 1981) which utilized six one-liter tanks; one control and five treatments with a dilution ratio of 0.65 between each treatment. The flow to each tank was 10 ml/min giving 14.4 volume additions per day. One control and five dilutions at 100, 80, 60, 40 and 20 percent of the stock concentration comprised the exposure series. The test solution</p>

depth was measured at 4-5 cm. No replicates were used. Tanks were placed in a random sequence in a constant temperature water bath. The water temperature was 25.4 deg C. The pH was 7.4. The dissolved oxygen was 6.9 mg/L. LC50 = 720 ug/L 95% C.L. (618-839 ug/L) and

Endpoint value

Mortality

Hour	Concentration	A	B	C	D	E
Initial	10	10	10	10	10	10
24			1	9	10	10
48			1	9	10	10
72			1	9	10	10
96			1	9	10	10

EC50=688 ug/L 95% C.L.(606-782 ug/L).

Mortality

Hour	Concentration	A	B	C	D	E
Initial	10	10	10	10	10	10
24			1	10	10	10
48			10	10	10	10
72			10	10	10	10
96			1	10	10	10

Measured concentrations as mg/L 1.81 mg/L

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Guideline study.

Reference Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.

Substance Name	Terpinolene
CAS No.	586-62-9
Remarks for Substance	Terpinolene in neat form had purity of >99%, in toxicant saturated water of 75% purity, in exposure water 41% purity
Method/guideline	Calculated LC50 and EC50 using the Trimmed Spearman-Kärber Method [Hamilton <i>et al.</i> , 1977] and corrected average of the analyzed tank concentrations
Test Type	Experimental
GLP	Ambiguous
Year	1990
Species/Strain/Supplier	Fathead minnows/American Tokyo Kasei
Exposure Period	96 hour
Analytical monitoring	GC Analysis
Remarks for Test Conditions	96 hour LC50 and EC50 tests were performed with fathead minnows using terpinolene. Test protocol was a continuous flow-through system. Tests were conducted in the electronic

diluter using 0.2 L test chambers.

The ecotoxicity test in fish was conducted with a modified Benoit diluter (Benoit, 1981) which utilized six one-liter tanks; one control and five treatments with a dilution ratio of 0.65 between each treatment. The flow to each tank was 10 ml/min giving 14.4 volume additions per day. One control and five dilutions at 100, 80, 60, 40 and 20 percent of the stock concentration comprised the exposure series. The test solution depth was measured at 4-5 cm. No replicates were used. Tanks were placed in a random sequence in a constant temperature water bath. The water temperature was 25 deg C. The pH was 7.6. The dissolved oxygen was 6.3 mg/L. Hardness was 45.1 mg/L CaCO₃ with an alkalinity of 42.6 mg/L CaCO₃.

Endpoint value

LC50 = 1210 ug/L 95% C.L. (1110-1330 ug/L)

Mortality

Hour	Concentration	A	B	C	D	E
Initial	9	10	10	10	10	10
24					5	9
48					6	10
72					7	10
96					7	10

EC50=763 ug/L.

Mortality

Hour	Concentration	A	B	C	D	E
Initial	9	10	10	10	10	10
24					7	10
48				10	10	10
72				0	10	10
96				10	10	10

Measured concentrations as mg/L

3.38 mg/L

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Guideline study.

Reference

Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Fish
Exposure Period	96 hour

Remarks for Test Conditions Input parameters: Melting point, -74.35, Water solubility - 13.8 mg/L, Log Kow - 5.3
Endpoint value LC50 = 0.221 mg/L
Data Qualities Reliabilities Reliability code 4. Not assignable.
Remarks for Data Reliability Code 4. Calculated.
Reference ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	138-86-3
CAS No.	<i>d</i> -Limonene (dipentene)
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Fish
Exposure Period	96 hour
Remarks for Test Conditions	Input parameters: Melting point, -74.35, Water solubility - 13.8 mg/L, Log Kow - 5.3
Endpoint value	LC50 = 0.221 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Fish
Exposure Period	96 hour
Remarks for Test Conditions	Input parameters: Melting point, -29.50, Water solubility - 9.5 mg/L, Log Kow - 4.88
Endpoint value	LC50 = 0.198 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Fish
Exposure Period	96 hour
Remarks for Test Conditions	Input parameters: Melting point, -64.83, Water solubility - 5.6mg/L, Log Kow - 4.8
Endpoint value	LC50 = 0.198 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Fish
Exposure Period	96 hour
Remarks for Test Conditions	Input parameters: Melting point, -66.11, Log Kow - 4.88
Endpoint value	LC50 = 0.201 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

3.2 Acute Toxicity to Aquatic Invertebrates

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5

Remarks for Substance	<i>d</i> -Limonene in neat form had purity of 99%, in toxicant saturated water of 85%, in exposure water 67%																												
Method/guideline	Flow-through. Calculated LC50 and EC50 using the Trimmed Spearman-Kärber Method [Hamilton <i>et al.</i> , 1977] and corrected average of the analyzed tank concentrations																												
Test Type	Experimental																												
GLP	No																												
Species/Strain/Supplier	<i>Daphnia Magna</i>																												
Analytical procedures	GC Analysis																												
Test Details	48 hours																												
Remarks for Test Conditions	<p>48 hour LC50 and EC50 tests were performed with daphnia magna from two different suppliers. Test protocol was a continuous flow-through system.</p> <p>The daphnia test was conducted with a modified Benoit diluter (Benoit, 1981) which utilized six one-liter tanks; one control and five treatments with a dilution ratio of 0.65 between each treatment. The flow to each tank was 10 ml/min giving 14.4 volume additions per day. One control and five dilutions at 100, 80, 60, 40 and 20 percent of the stock concentration comprised the exposure series. The test solution depth was measured at 4-5 cm. No replicates were used. Stainless steel screen enclosures (forty mesh) were placed in each chamber for the daphnia exposures. Tanks were placed in a random sequence in a constant temperature water bath.</p> <p>A saturator flow rate of 21 mL/min resulted in exposure concentrations of 251 to 1890 ug/L.</p> <p>The water temperature was 20 deg C. The pH was 7.9. The dissolved oxygen was 7.8 mg/L.</p>																												
Measured concentrations as mg/L	2.73 mg/L																												
EC50, EL50, LC0, at 24,48 hours	48 hr LC50 = .577 mg/L or 577 ug/L 95% C.L. (496-672 ug/L)																												
	<table border="0"> <tr> <td colspan="7">Mortality</td> </tr> <tr> <td>Hour</td> <td>Concentration</td> <td>A</td> <td>B</td> <td>C</td> <td>D</td> <td>E</td> </tr> <tr> <td>Initial</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> </tr> <tr> <td>48</td> <td></td> <td>7</td> <td>15</td> <td>15</td> <td>15</td> <td></td> </tr> </table>	Mortality							Hour	Concentration	A	B	C	D	E	Initial	15	15	15	15	15	15	48		7	15	15	15	
Mortality																													
Hour	Concentration	A	B	C	D	E																							
Initial	15	15	15	15	15	15																							
48		7	15	15	15																								
	EC50=.421mg/L or 421 ug/L (no reliable confidence limits)																												
	<table border="0"> <tr> <td colspan="7">Mortality</td> </tr> <tr> <td>Hour</td> <td>Concentration</td> <td>A</td> <td>B</td> <td>C</td> <td>D</td> <td>E</td> </tr> <tr> <td>Initial</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> </tr> <tr> <td>48</td> <td></td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> <td>15</td> </tr> </table>	Mortality							Hour	Concentration	A	B	C	D	E	Initial	15	15	15	15	15	15	48		15	15	15	15	15
Mortality																													
Hour	Concentration	A	B	C	D	E																							
Initial	15	15	15	15	15	15																							
48		15	15	15	15	15																							
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.																												
Data Reliability Remarks	Code 1. Guideline study.																												
Reference	Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.																												

Substance Name	<i>d</i> -Limonene						
CAS No.	5989-27-5						
Remarks for Substance	<i>d</i> -Limonene in neat form had purity of 99%, in toxicant saturated water of 85%, in exposure water 67%						
Method/guideline	Calculated LC50 and EC50 using the Trimmed Spearman-Kärber Method [Hamilton <i>et al.</i> , 1977] and corrected average of the analyzed tank concentrations						
Test Type	Experimental						
GLP	No						
Species/Strain/Supplier	<i>Daphnia Magna</i>						
Analytical procedures	GC Analysis						
Test Details	48 hours						
Remarks for Test Conditions	<p>48 hour LC50 and EC50 tests were performed with daphnia magna using <i>d</i>-limonene from two different suppliers. Test protocol was a continuous flow-through system.</p> <p>The daphnia test was conducted with a modified Benoit diluter (Benoit, 1981) which utilized six one-liter tanks; one control and five treatments with a dilution ratio of 0.65 between each treatment. The flow to each tank was 10 ml/min giving 14.4 volume additions per day. One control and five dilutions at 100, 80, 60, 40 and 20 percent of the stock concentration comprised the exposure series. The test solution depth was measured at 4-5 cm. No replicates were used. Stainless steel screen enclosures (forty mesh) were placed in each chamber for the daphnia exposures. Tanks were placed in a random sequence in a constant temperature water bath. A saturator flow rate of 21 mL/min resulted in exposure concentrations of 178 to 1350 ug/L.</p> <p>The water temperature was 19.7 deg C. The pH was 7.9. The dissolved oxygen was 8.0 mg/L. Hardness was 45.5 mg/L CaCO₃ with an alkalinity of 36.6 mg/L CaCO₃.</p>						
Measured concentrations as mg/L	1.81 mg/L						
EC50, EL50, LC0, at 24,48 hours	48 hr LC50 = .924 mg/L (no reliable confidence limits)						
	Mortality						
	Hour	Concentration	A	B	C	D	E
	Initial	15	15	15	15	15	15
	48		15	15	15	15	15
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.						
Data Reliability Remarks	Code 1. Guideline study.						
Reference	Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.						

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	97% purity
Method/guideline	Static. EC50
Test Type	Experimental
GLP	Ambiguous
Year	1987
Species/Strain/Supplier	<i>Daphnia pulex</i>
Analytical procedures	GC Analysis
Test Details	48 hours
Remarks for Test Conditions	The 48 hour static acute tests with <i>Daphnia pulex</i> were conducted without feeding the organisms. Tests included a water or solvent control and five concentrations of toxicant dissolved in water or toxicant. Tests chambers contained at least 10 neonates in 150 ml of water. Two replicates were performed.
EC50, EL50, LC0, at 24,48 hours	EC50 = 69.6 mg/L at 48 hour
Data Qualities Reliabilities	Reliability code 3. Not reliable.
Data Reliability Remarks	Code 3. Test concentration exceeds the limit of solubility of <i>d</i> -limonene.
Reference	Passino D.R., and Smith S. (1987) Acute bioassays and hazard evaluation of representative contaminants detected in great lakes fish. <i>Environmental Toxicology and Chemistry</i> , 6, 901.

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	Flow-through. Calculated LC50 and EC50 using the Trimmed Spearman-Kärber Method [Hamilton <i>et al.</i> , 1977] and corrected average of the analyzed tank concentrations
Test Type	Experimental
GLP	No
Species/Strain/Supplier	<i>Daphnia Magna</i>
Analytical procedures	GC Analysis
Test Details	48 hours
Remarks for Test Conditions	48 hour LC50 and EC50 tests were performed with <i>daphnia magna</i> using terpinolene. Test protocol was a continuous flow-

through system.

The daphnia test was conducted with a modified Benoit diluter (Benoit, 1981) which utilized six one-liter tanks; one control and five treatments with a dilution ratio of 0.65 between each treatment. The flow to each tank was 10 ml/min giving 14.4 volume additions per day. One control and five dilutions at 100, 80, 60, 40 and 20 percent of the stock concentration comprised the exposure series. The test solution depth was measured at 4-5 cm. No replicates were used. Stainless steel screen enclosures (forty mesh) were placed in each chamber for the daphnia exposures. Tanks were placed in a random sequence in a constant temperature water bath.

The water temperature was 19.6 deg C. The pH was 7.7. The dissolved oxygen was 6.2 mg/L.

A saturator flow rate of 21 mL/min resulted in exposure concentrations of 279 to 3020 ug/L.

Measured concentrations as mg/L 3.38 mg/L
EC50, EL50, LC0, at 24,48 hours 48 hr LC50 = 2.55 mg/L or 2550 ug/L (no reliable confidence limits)

Mortality
Hour Concentration A B C D E
Initial 15 15 15 15
48 13

and EC50=1380 ug/L (no reliable confidence limits)

Mortality
Hour Concentration A B C D E
Initial 15 15 15 15
48 15 15

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Data Reliability Remarks Code 1. Guideline study.

Reference Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	EPA Committee on Methods for Toxicity Tests with Aquatic Organisms (without replicate concentrations)
Test Type	Experimental
GLP	Ambiguous
Year	1986

Species/Strain/Supplier	<i>Daphnia Magna</i>
Test Details	48 hours
Remarks for Test Conditions	200 ml of the test solution and 10 Daphnia were used. Dissolved oxygen and pH were determined initially and at 48 hour. Mortalities were recorded at 24 and 48 hour.
EC50, EL50, LC0, at 24,48 hours	LC50 = 31 mg/L at 48 hour
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Data Reliability Remarks	Code 2. Basic data given: comparable to guidelines/standards.
Reference	Waggy G.L. and Blessing R.L. (1986) Private communication.

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	Brine shrimp lethality test [Meyer, 1982]
Test Type	Experimental
GLP	Ambiguous
Year	1998
Species/Strain/Supplier	Brine shrimp
Analytical procedures	GC Analysis
Test Details	24 hours
Remarks for Test Conditions	<p>The essential oil of <i>Apium graveolens</i> (celery) was analyzed by GC-MS. Components were isolated using dry column chromatography and low pressure liquid chromatography to isolate the hydrocarbon component and individual hydrocarbon compounds, respectively. The brine shrimp lethality test was performed using the crude volatile oil, the hydrocarbon and oxygenated fractions, and the isolated compounds. The results given here are for the respective isolated compounds as listed under Substance Name.</p> <p>Brine shrimp eggs were hatched in seawater and used after 48 hrs. Ten shrimps were added to three vials for each of the following doses (2, 50 and 200 ppm). The vials were prepared by thoroughly mixing with the allotted amount of artificial seawater to achieve the correct concentration. The number of deaths out of 30 shrimps per dose were recorded after 24 hrs. LC50 values and corresponding 95% confidence intervals were determined. Compounds with LC50 values over 200 ppm were retested at 1000 ppm. LC50 values greater than 200 ppm were considered inactive for pure compounds.</p>
EC50, EL50, LC0, at 24,48 hours	LC50 = 39.2 mg/L

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Data Reliability Remarks Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.

Reference Saleh M.M., Hashem F.A., Glombitza K.A. (1998) Cytotoxicity and in vitro effects on human cancer cell lines of volatiles of *Apium graveolens* var. *filicinum*. *Pharmaceutical Pharmacological Letters*, 8(2), 97-99.

Substance Name	Dihydromyrcene
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CAS No. 2436-90-0

Method/guideline Brine shrimp lethality test [Meyer, 1982]

Test Type Experimental

GLP Ambiguous

Year 1998

Species/Strain/Supplier Brine shrimp

Analytical procedures GC Analysis

Test Details 24 hours

Remarks for Test Conditions The essential oil of *Apium graveolens* (celery) was analyzed by GC-MS. Components were isolated using dry column chromatography and low pressure liquid chromatography to isolate the hydrocarbon component and individual hydrocarbon compounds, respectively. The brine shrimp lethality test was performed using the crude volatile oil, the hydrocarbon and oxygenated fractions, and the isolated compounds. The results given here are for the respective isolated compounds as listed under Substance Name.

Brine shrimp eggs were hatched in seawater and used after 48 hrs. Ten shrimps were added to three vials for each of the following doses (2, 50 and 200 ppm). The vials were prepared by thoroughly mixing with the allotted amount of artificial seawater to achieve the correct concentration. The number of deaths out of 30 shrimps per dose were recorded after 24 hrs. LC50 values and corresponding 95% confidence intervals were determined. Compounds with LC50 values over 200 ppm were retested at 1000 ppm. LC50 values greater than 200 ppm were considered inactive for pure compounds.

EC50, EL50, LC0, at 24,48 hours LC50 =104.1 mg/L

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Data Reliability Remarks Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.

Reference Saleh M.M., Hashem F.A., Glombitza K.A. (1998) Cytotoxicity and invitro effects on human cancer cell lines of volatiles of *Apium graveolens* var. *filicinum*. *Pharmaceutical*

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	<i>Daphnia Magna</i>
Test Details	48 hours
EC50, EL50, LC0, at 24,48 hours	LC50 = 0.496 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Data Reliability Remarks	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	<i>Daphnia Magna</i>
Test Details	48 hours
EC50, EL50, LC0, at 24,48 hours	LC50 = 0.612 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Data Reliability Remarks	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	<i>Daphnia Magna</i>

Test Details	48 hours
EC50, EL50, LC0, at 24,48 hours	LC50 = 1.147 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Data Reliability Remarks	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	<i>Daphnia Magna</i>
Test Details	48 hours
EC50, EL50, LC0, at 24,48 hours	LC50 = 0.263 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Data Reliability Remarks	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

3.3 Acute Toxicity to Aquatic Plants

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	<i>d</i> -Limonene in neat form had purity of 99%, in toxicant saturated water of 85%, in exposure water 67%
Method/guideline	ASTM methods (ASTM, 1988)
Test Type	Static 96 hour toxicity test with micro algae
GLP	Ambiguous
Year	1990
Species/Strain/Supplier	Green algae/ <i>Selenastrum capricornutum</i>
Exposure Period	96 hour
Analytical monitoring	GC analysis

Remarks for Test Conditions	Test conditions followed ASTM methods. Changes were made to help reduce any problems associated with volatility. GC analysis was performed at 0, 24, and 96 hours. The 96 hour sample was performed only if the chemical was detected at the 24 hour sampling. Test cell concentrations were approximately 10000 cells/mL. Coulter counters and electronic particle counter were used to count cells and determine mean cell volume. The IC50 (the concentration at which 50% growth inhibition) was calculated using a linear interpolation program.
Nominal concentrations as mg/L	Not given
Measured concentrations as mg/L	1.81 mg/L
Endpoint value	No significant inhibition
Conclusion Remarks	No significant inhibition.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
Reference	Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.

Substance Name	Terpinolene
CAS No.	586-62-9
Remarks for Substance	Terpinolene in neat form had purity of >99%, in toxicant saturated water of 75% purity, in exposure water 41% purity
Method/guideline	ASTM methods (ASTM, 1988)
Test Type	Static 96 hour toxicity test with micro algae
GLP	Ambiguous
Year	1990
Species/Strain/Supplier	Green algae/ <i>Selenastrum capricornutum</i>
Exposure Period	96 hour
Analytical monitoring	GC analysis
Remarks for Test Conditions	Test conditions followed ASTM methods. Changes were made to help reduce any problems associated with volatility. GC analysis was performed at 0, 24, and 96 hours. The 96 hour sample was performed only if the chemical was detected at the 24 hour sampling. Test cell concentrations were approximately 10000 cells/mL. Coulter counters and electronic particle counter were used to count cells and determine mean cell volume. The IC50 (the concentration at which 50% growth inhibition) was calculated using a linear interpolation program.
Nominal concentrations as mg/L	Not given

Measured concentrations as mg/L	3.38 mg/L
Endpoint value	No significant inhibition
Conclusion Remarks	No significant inhibition.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
Reference	Broderius S. (1990) Toxicity of Eight Terpenes to Fathead Minnows, Daphnids, and Algae. U.S. EPA Environmental Research Laboratory-Duluth and ASCI Corporation.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Green algae
Exposure Period	96 hours
Endpoint value	EC50 = 0.360 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Myrcene
CAS No.	123-35-3
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Green algae
Exposure Period	96 hours
Endpoint value	EC50 = 0.813 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Terpinolene
CAS No.	586-62-9
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Green algae
Exposure Period	96 hours
Endpoint value	EC50 = 0.441 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Method/guideline	ECOSAR
Test Type	Calculated
Species/Strain/Supplier	Green algae
Exposure Period	96 hours
Endpoint value	EC50 = 0.194 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4. Calculated.
Reference	ECOSAR EPI Suite (2000) U.S. Environmental Protection Agency (Nabholz V. and G. Cash, 1998).

4 Human Health Toxicity

4.1 Acute Toxicity

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Test Substance	Purity not given
Method/guideline	Litchfield and Wilcoxon, 1949
Test Type	Oral LD 50
GLP	No
Year	1975
Species/strain	Rat/Wistar
Sex	Male and Female
# of animals per sex per dose	10 male 10 female
Vehicle	Arabic gum/water
Route of Administration	Oral
Remarks for Test Conditions	Ten male and ten female rats per group were used.
Value LD50 or LC50 with confidence limits	Male LD50 = 4400 (3400-5900) mg/kg bw (95% confidence limit) and female LD50= 5.2 (3.9-7.0) g/kg bw (95% confidence limit)
Number of deaths at each dose level	1500 mg/kg NOE in males. Not tested in females. 1900 mg/kg LTL 30% mortality in males. No observed effects in females. 2500 mg/kg LTL 30% mortality in males. 20% mortality in females. 3.3 g/kg LTL 40% mortality in males. 50% mortality in females. 4.3 g/kg LTL 60% mortality in males. 40% mortality in females. 4.4 g/kg LD50 male rats. 95% Confidence limits (3.4-5.9). 5.2 g/kg LD50 female rats. 95% Confidence limits (3.9-7.0). 5.6 g/kg LTL 40% mortality in males. 50% mortality in females. 7.3 g/kg LTL 90% mortality in males. 60% mortality in females. 9.4 g/kg LTL 70% mortality in males. 60% mortality in females. 12.2 g/kg LTL 80% mortality in males. 90% mortality in females. 15.9 g/kg LTL 90% mortality in males. 100% mortality in females.
Remarks for Results	The male LD50 = 4400 (3400-5900) mg/kg bw (95% confidence limits) and female LD50= 5200 (3900-7000) mg/kg bw (95% confidence limits).
Conclusion remarks	The male LD50 = 4400 (3400-5900) mg/kg bw (95% confidence limits) and female LD50= 5200 (3900-7000) mg/kg bw (95% confidence limits).
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.

Remarks for Data Reliability Code 2. Basic data given: comparable to guidelines/standards.

References Tsuji M., Y.Fujisaki, Y.Arikawa, S.Masuda, S.Kinoshita, A.Okubo, K.Noda, H.Ide and Y.Iwanaga (1975a) Studies on *d*-limonene as a gallstone solubilizer: Acute and sub-acute toxicities. Journal Oyo Yakuri, 9, 387-401.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Test Substance	Purity not given
Method/guideline	Litchfield and Wilcoxon, 1949
Test Type	Oral LD 50
GLP	No
Year	1975
Species/strain	Mouse
Sex	Male and Female
# of animals per sex per dose	10 male 10 female
Vehicle	Arabic gum/water
Route of Administration	Oral
Remarks for Test Conditions	Ten male and ten female mice per group were used.
Value LD50 or LC50 with confidence limits	Male LD50 = 5600 (4800-6500) mg/kg bw (95% confidence limit) and female LD50 = 6600 (5500-7900) mg/kg bw (95% confidence limit)
Number of deaths at each dose level	3.0 g/kg NOE in male mice. 3.5 g/kg LTL 10% mortality in male mice. No observable effects in female mice. 4.3 g/kg LTL 20% mortality in female mice. No observable effects in male mice. 5.3 g/kg LTL 20% mortality in female mice. 30% mortality in male mice. 5.6 g/kg LD50 male mice. 95% confidence limits (4.8-6.5) 6.6 g/kg LD50 female mice. 95% confidence limits (5.5-7.9). 7.0 g/kg LTL 30% mortality in female mice. 60% mortality in male mice. 7.5 g/kg LTL 60% mortality in female mice. 90% mortality in male mice. 8.3 g/kg LTL 90% mortality in female mice. 100% mortality in male mice. 10.0 g/kg LTL 100% mortality in female mice.
Remarks for Results	The male LD50 = 5600 (4800-6500) mg/kg bw (95% confidence limits) and female LD50 = 6600 (5500-7900) mg/kg bw (95% confidence limits).
Conclusion remarks	Male LD50 = 4400 (3400-5900) mg/kg bw (95% confidence limit) and female LD50 = 5200 (3900-7000) mg/kg bw (95% confidence limits).
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.

References

Tsuji M., Y.Fujisaki, Y.Arikawa, S.Masuda, S.Kinoshita, A.Okubo, K.Noda, H.Ide and Y.Iwanaga (1975a) Studies on *d*-limonene as a gallstone solubilizer: Acute and sub-acute toxicities. Journal Oyo Yakuri, 9, 387-401.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Oral LD 50
GLP	No
Year	1972
Species/strain	Rat/Wistar
Sex	Male
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Oral
Remarks for Test Conditions	Ten male albino Wistar rats per group were used. Animals were fasted for a minimum of 16 hours prior to administration of the test material. Animals weighed 200-250 grams. Following dosing the animals received food and water ad libitum. Observations for mortality were made at 1 and 6 hours after dosing and daily thereafter for 14 days. Toxic effects were also observed. Gross necropsies were performed on all survivors.
Value LD50 or LC50 with confidence limits	Greater than 5000 mg/kg bw
Number of deaths at each dose level	None
Remarks for Results	Animals experienced lethargy. No deaths occurred. Oral LD50 greater than 5000 mg/kg.
Conclusion remarks	Oral LD50 greater than 5000 mg/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1972c) Acute oral toxicity of limonene in rats. Unpublished report.

Substance Name	Terpinolene
CAS No.	586-62-9

Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Oral LD 50
GLP	No
Year	1975
Species/strain	Rat
Sex	Not reported
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Oral
Value LD50 or LC50 with confidence limits	4.39 ml/kg 95% Confidence Limits (3.75-5.14 ml/kg)
Number of deaths at each dose level	3.0 ml/kg 0 deaths 3.5 ml/kg 1 death 4.0 ml/kg 5 deaths 5.0 ml/kg 6 deaths
Remarks for Results	The oral LD50 was calculated to be 4.39 ml/kg bw with 95% C.L. (3.75-5.14 ml/kg).
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Levenstein I. (1975) Oral LD 50 of terpinolene in rats. Unpublished report to RIFM.

Substance Name	Myrcene
Remarks for Test Substance	Purity not given
CAS No.	123-35-3
Method/guideline	Not given
Test Type	Oral LD 50
GLP	Ambiguous
Year	1990
Species/strain	Rat/Wistar
Sex	Male and Female
# of animals per sex per dose	1 male 1 female except three highest doses
Vehicle	Corn oil
Route of Administration	Oral-Gavage

Remarks for Test Conditions	Two animals (1 male and 1 female) were used per dose with the exception of the three highest doses where 2 animals of each sex were used (see number of deaths at each dose level). Animals were fasted overnight prior to test administration. Animals were observed for 14 days following administration. Necropsies were performed on all animals.
Value LD50 or LC50 with confidence limits	Greater than 11.39 g/kg bw
Number of deaths at each dose level	0 g/kg bw Male 0/1 Female 0/1 0.67 g/kg bw Undefined results 1.00 g/kg bw Male 0/1 Female 0/1 1.5 g/kg bw Male 0/1 Female 0/1, 2.25 g/kg bw Male 0/1 Female 0/1 3.25 g/kg bw Male 0/1 Female 0/1 5.06 g/kg bw Male 0/2 Female 0/2 7.59 g/kg bw Male 0/2 Female 0/2 11.39 g/kg bw Male 0/2 Female 0/2
Remarks for Results	Animals at the three highest dose levels experienced palpebral ptosis, hypoactivity, and ataxia.
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Paumgarten F.J.R. (1990) Single Dose Toxicity of <i>beta</i> -myrcene; a natural analgesic substance. Brazilian J Med Biol Res 23; 873-877.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Oral LD 50
GLP	Ambiguous
Year	1990
Species/strain	Mouse/Albino Swiss
Sex	Male and Female
# of animals per sex per dose	1 male 1 female with the exception of 2 doses
Vehicle	Corn oil
Route of Administration	Oral-Gavage
Remarks for Test Conditions	Two animals (1 male and 1 female) were used per dose with the exception of two doses where 3 animals of each sex were used (see number of deaths at each dose level). Animals were fasted overnight prior to test administration. Animals were observed for 14 days following administration. Necropsies were performed on all animals.
Value LD50 or LC50 with confidence limits	5060 mg/kg bw
Number of deaths at each	0 g/kg bw Male 0/1 Female 0/1 0.67 g/kg bw Undefined results

dose level	1.00 g/kg bw Male 0/1 Female 0/1 1.5 g/kg bw Male 0/1 Female 0/1
	2.25 g/kg bw Male 0/1 Female 0/1
	3.25 g/kg bw Male 0/1 Female 0/1
	5.06 g/kg bw Male 2/3 Female 3/3
	7.59 g/kg bw Male 3/3 Female 2/3
	11.39 g/kg bw Male 1/1 Female 1/1
Remarks for Results	Animals at the three highest dose levels experienced palpebral ptosis, hypoactivity, and ataxia.
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Paumgarten F.J.R. (1990) Single Dose Toxicity of <i>beta</i> -myrcene; a natural analgesic substance. Brazilian J Med Biol Res 23; 873-877.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Oral LD 50
GLP	No
Year	1972
Species/strain	Rat/Wistar
Sex	Male
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Oral

Remarks for Test Conditions	Ten male albino Wistar rats per group were used. Animals were fasted for a minimum of 16 hours prior to administration of the test material. Animals weighed 200-250 grams. Following dosing the animals received food and water ad libitum. Observations for mortality were made at 1 and 6 hours after dosing and daily thereafter for 14 days. Toxic effects were also observed. Gross necropsies were performed on all survivors. Greater than 5000 mg/kg bw
Value LD50 or LC50 with confidence limits	
Number of deaths at each dose level	1/10 deaths. Death occurred overnight following administration. Clinical signs included lethargy and urinary incontinence.
Remarks for Results	Animals experienced lethargy urinary incontinence. One deaths occurred. Oral LD50 greater than 5000 mg/kg.
Conclusion remarks	Oral LD50 greater than 5000 mg/kg.
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1972d) Acute oral toxicity of myrcene in rats. Unpublished report.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Remarks for Test Substance	Purity not given
Method/guideline	Litchfield and Wilcoxon, 1949
Remarks for Substance	Clear liquid
Test Type	Oral LD 50
GLP	No
Year	1980
Species/strain	Rat/Wistar
Sex	Male
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Oral
Remarks for Test Conditions	The rats were observed for 3-4 hours after dosing and once daily for 14 days. Mortality, toxicology, and pharmacological effects were recorded.
Value LD50 or LC50 with confidence limits	Greater than 5000 mg/kg bw
Number of deaths at each dose level	1 death at 5000 mg/kg bw
Remarks for Results	Lethargy and piloerection were noted 3-4 hours post dose. Most animals were generally healthy thereafter.

Conclusion remarks	Oral LD50 greater than 5000 mg/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1980b) Acute oral toxicity of dihydromyrcene in rats. Unpublished report.

Substance Name	Orange peel oil, sweet (Citrus sinensis (L.) Osbeck)
CAS No.	8008-57-9
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Oral LD 50
GLP	No
Year	1973
Species/strain	Rat/Wistar
Sex	Male
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Oral
Remarks for Test Conditions	The rats were observed for 3-4 hours after dosing and once daily for 14 days. Mortality, toxicology, and pharmacological effects were recorded.
Value LD50 or LC50 with confidence limits	Greater than 5000 mg/kg bw
Remarks for Results	The oral LD50 was determined to be greater than 5000 mg/kg bw.
Conclusion remarks	Oral LD50 >5000 mg/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1973b) Acute oral toxicity of orange oil in rats. Unpublished report.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Test Substance	Purity not given

Method/guideline	Not given
Remarks for Substance	Clear liquid
Test Type	Dermal LD50
GLP	No
Year	1972
Species/strain	Rabbit/New Zealand White
Sex	Not reported
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Dermal
Remarks for Test Conditions	A single 24 hour application of limonene (5 g/kg) was applied to the clipped abraded abdominal skin of 10 rabbits weighing from 1.9 to 2.4 kg. Observations for mortality and toxic effects were made for seven days following exposure. Gross necropsies were performed on all animals at study termination.
Value LD50 or LC50 with confidence limits	Greater than 5000 mg/kg bw
Number of deaths at each dose level	None
Remarks for Results	No animals died during the study. Additionally, there was no evidence of toxicity resulting from application of the material.
Conclusion remarks	Oral LD50 greater than 5000 mg/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1972a) Acute dermal toxicity of limonene in rabbits. Unpublished report.

Substance Name	Dihydromyrcene
CAS No.	2436-90-0
Remarks for Test Substance	Purity not given
Method/guideline	Litchfield and Wilcoxon, 1949
Remarks for Substance	Clear liquid
Test Type	Dermal LD50
GLP	No
Year	1980

Species/strain	Rabbit/New Zealand White
Sex	Male
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Dermal
Remarks for Test Conditions	A single 24 hour application of dihydromyrcene (5 g/kg) was applied to the clipped abraded abdominal skin of 10 rabbits. Observations for mortality and toxic effects were made for 14 days following exposure. All animals were examined for gross pathology.
Value LD50 or LC50 with confidence limits	Greater than 5000 mg/kg bw
Number of deaths at each dose level	1 death at 5000 mg/kg bw
Remarks for Results	Toxic signs noted infrequently included lethargy and diarrhea. Internal organs of surviving animals were normal following superficial examination. One animal showed signs of cardiac and respiratory abnormalities. Most also had skin abnormalities.
Conclusion remarks	Dermal LD50 >5000 mg/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1980b) Acute dermal toxicity of dihydromyrcene in rabbits. Unpublished report.

Substance Name	Terpinolene
CAS No.	586-62-9
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Dermal LD50
GLP	No
Year	1975
Species/strain	Rat
Sex	Not reported
# of animals per sex per dose	4
Vehicle	None
Route of Administration	Dermal

Value LD50 or LC50 with confidence limits	Greater than 5 ml/kg bw
Number of deaths at each dose level	0 deaths
Remarks for Results	The dermal LD50 was determined to be greater than 5 ml/kg bw.
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Levenstein I. (1975) Dermal LD 50 of terpinolene in rats. Unpublished report to RIFM.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Dermal LD50
GLP	No
Year	1972
Species/strain	Rabbit/New Zealand White
Sex	Not reported
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Dermal
Remarks for Test Conditions	A single 24 hour application of terpinolene (5 g/kg) was applied to the clipped abraded abdominal skin of 10 rabbits weighing from 1.9 to 2.4 kg. Observations for mortality and toxic effects were made for seven days following exposure. Gross necropsies were performed on all animals at study termination.
Value LD50 or LC50 with confidence limits	Greater than 5000 mg/kg bw
Number of deaths at each dose level	No deaths during the course of the study. No evidence of toxicity from percutaneous absorption of the test substance. Erythema and edema were reported during the first few days of observation, but cleared by the study termination.
Remarks for Results	The dermal LD50 was determined to be greater than 5000 mg/kg bw.
Conclusion remarks	Dermal LD50 greater than 5000 mg/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.

References Moreno O. (1972b) Acute dermal toxicity of myrcene in rabbits. Unpublished report.

Substance Name	Orange peel oil, sweet (Citrus sinensis (L.) Osbeck)
CAS No.	123-35-3
Remarks for Test Substance	Purity not given
Method/guideline	Not given
Test Type	Dermal LD50
GLP	No
Year	1973
Species/strain	Rabbit/New Zealand White
Sex	Male
# of animals per sex per dose	10
Vehicle	None
Route of Administration	Dermal
Remarks for Test Conditions	A single 24 hour application of terpinolene (5 g/kg) was applied to the clipped abraded abdominal skin of 10 rabbits weighing from 1.9 to 2.4 kg. Observations for mortality and toxic effects were made for seven days following exposure. Gross necropsies were performed on all animals at study termination. Greater than 5000 mg/kg bw
Value LD50 or LC50 with confidence limits	
Remarks for Results	The dermal LD50 was determined to be greater than 5000 mg/kg bw.
Conclusion remarks	Dermal LD50 greater than 5.0 g/kg bw
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Moreno O. (1973a) Acute dermal toxicity of orange oil in rabbits. Unpublished report.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Test Substance	Purity not given
Remarks for Substance	Purity undetermined
Method/guideline	Not given

Test Type	Inhalation ED25
GLP	Not reported
Year	1977
Species/strain	Mouse
Sex	Female
# of animals per sex per dose	5
Vehicle	None
Route of Administration	Inhalation
Remarks for Test Conditions	The respiratory irritation potential of fragrance raw materials was assessed in CF-1 females by recording respiratory rate using a whole body plethysmograph. Mice were exposed to test materials for 1 min using a nebulizer for aerosolization in a 2600 ml chamber. Materials shown to be sensory irritants were further tested in mice cannulated via the trachea & compared to an intact mouse breathing through its nose. Comparisons made were between the pre-exposure & exposure rate values for each material at each dose level. Materials were of undetermined purity. Respiratory tract, via nose, mild-moderate resp depression; ED25 = 570 ug/l; no effects when inhaled through tracheal cannula. (Troy, 1977) 9011
Value LD50 or LC50 with confidence limits	No ED25 determined.
Number of deaths at each dose level	None
Remarks for Results	Slight respiratory depression. Lower tract exposures not performed. No dose-response relationship.
Data Qualities Reliabilities	Reliability code 2. Reliable with restrictions.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Troy W.R. (1977) Doctoral Dissertation: The comparative respiratory irritation potential of fourteen fragrance raw materials. Unpublished report to RIFM.

4.2 Genetic Toxicity

4.2.1 In vitro Genotoxicity

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Ames test
Test Type	Ames reverse mutation

System of Testing	Bacterial
GLP	No
Year	1980
Species/Strain	<i>Salmonella typhimurium</i> TA 98, TA 100, TA 1535 and TA 1537
Metabolic Activation	With and without rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	0.03, 0.3, 3 and 30 micromol/plate for TA98 and TA100; 3 micromol/plate for the remaining strains
Remarks for Test Conditions	The solvent used was ethanol. Only one replicate was performed for the substances, which tested negative. Initially a Difco nutrient broth was used, but was subsequently suspected of weak mutagenic activity. The broth was changed to Oxoid nutrient broth No. 2. Revertants were scored on glucosenminimal salts medium supplemented with 0.05 umol histidine and 0.05 umol biotin. Plates used for viable counts contained 10 umol histidine (and 0.05umol biotin). The viable count was determined; the number of spontaneous revertants was measured; the presence of the rfa-mutation was checked by crystal violet inhibition; the presence of plasmid pKM 101 in strains TA 98 and TA100 was checked by resistance to ampicillin; the response to positive controls N-methyl-N'-nitro-N-nitrosoguanidin, and 2-aminoanthracene was checked.
Results	No mutagenic effects.
Cytotoxic concentration	Greater than 3 micromoles/plate
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Limonene was inactive in <i>Salmonella</i> strains TA 1535, TA 1537, TA 98 & TA 100 both in the presence and absence of metabolic activation system.
Conclusion Remarks	No evidence of mutagenic activity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Florin I., Rutberg L., Curvall M., and Enzell C.R. (1980) Screening of tobacco smoke constituents for mutagenicity using the Ames test. Toxicology, 18 pages 219-232.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Ames test –plate incorporation method
Test Type	Ames reverse mutation

System of Testing	Bacterial
GLP	No
Year	1989
Species/Strain	<i>Salmonella typhimurium</i> TA 98, TA 100, TA 1535, TA 1537, and TA1538
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	150000 micrograms/plate
Remarks for Test Conditions	The bacteria were cultured in Oxoid medium for approximately 12 hours. Selection of histidine revertants was performed in Vogel Bonner medium containing 2% glucose and 1.5% bacto agar. The preparation and composition of the overlay agar and a metabolic activation mix consisting of a 9000x g supernatant fraction (S9) of Aroclor 1254-induced adult male Sprague Dawley rat liver and necessary cofactors. The assays were conducted by addition of the test substance to a 2.0 ml volume of top agar along with a 0.1 ml aliquot of the bacterial culture and either the metabolic activation mix or an equivalent volume of phosphate buffer. These components were mixed rapidly and poured onto the minimal medium plates. After two days incubation at 37 °C, revertant colonies were counted.
Results	No mutagenic effects.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Limonene was inactive in Salmonella strains TA 1535, TA 1537, TA 1538, TA 98 & TA 100 both in the presence and absence of metabolic activation system.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Heck, J. D., Vollmuth, T. A., Cifone, M. A., Jagannath, D. R., Myhr B., and R.D. Curren (1989). An evaluation of food flavoring ingredients in a genetic toxicity screening battery The Toxicologist, 9(1), 257.

Substance Name	d-Limonene
CAS No.	5989-27-5
Remarks for Substance	Solvents used included DMSO and/or ethanol.
Method/guideline	Ames
Test Type	Ames reverse mutation

System of Testing	Bacterial
GLP	Ambiguous
Year	1993
Species/Strain	<i>Salmonella typhimurium</i> TA102
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats.
Doses/Concentration	Up to 5000 micrograms per plate
Statistical Methods	Not given
Remarks for Test Conditions	Bacteria were grown in a nutrient broth, and aliquots were frozen at -80 deg C. All experiments were performed from the same frozen batch. Limonene was tested in two independent experiments using five doses and three plates per dose. Solvents used for diluting the compounds were DMSO, ethanol and water. All compounds were tested up to 5000 micrograms per plate if possible unless limited by cytotoxicity or precipitation.
Results	Negative
Cytotoxic concentration	Not given
Genotoxic Effects	Negative
Appropriate statistical evaluations	Not given
Remarks for results	Limonene was inactive in <i>Salmonella</i> strains TA 102 in the presence and absence of metabolic activation system.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Muller W. (1993) Evaluation of mutagenicity testing with <i>Salmonella typhimurium</i> TA102 in three different laboratories. Environmental Health Perspectives Supplements. 101, 33-36.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	99.7% analyzed purity
Method/guideline	Ames test
Test Type	Ames reverse mutation
System of Testing	Bacterial
GLP	Ambiguous

Year	1983
Species/Strain	<i>Salmonella typhimurium</i> TA1535, TA 1537, TA98, TA100
Metabolic Activation	Rat liver microsomal fraction S9 from Aroclor induced Sprague Dawley rats
Doses/concentration levels	0.3-3333 micrograms/plate
Statistical Methods	Model used presented in Margolin <i>et al.</i> , 1981
Remarks for Test Conditions	<p>Cultures of each tester strain were prepared for storage as described in the Ames method paper. Frozen cultures were stored in liquid nitrogen (EGG) or in a -70 deg C freezer (CWR, SRI) in 0.2-ml aliquots or in 1-ml aliquots in sterile screw cap vials. To inoculate overnight cultures, one lab transferred a loopful of cells that were maintained on Columbia agar slants kept at 4 deg C into Columbia broth. The other lab (EGG) transferred a loopful of thawed cultures into Oxoid Nutrient broth #2 and discarded a unused portion of the thawed culture. SRI used all of the thawed 1-ml to inoculate minimal glucose medium. To check for toxicity, up to 10 mg/plate or the limit of solubility, both in the presence and absence of S-9 mix. One or more parameters were used as an indication of toxicity, viability on complete medium and reduced numbers of revertant colonies per plate and or thinning or absence of the bacterial lawn. If toxicity was not apparent, the highest dose tested was 10 mg/plate; otherwise the upper limit of solubility was used. If toxicity was observed the doses of the test substance were chosen so that the high dose exhibited some degree of toxicity. At least five doses of the test chemical, in addition to solvent and positive controls, were tested on each strain in the presence of S9 mix or buffer. Three plates were used and the experiment was repeated no less than 1 week after completion of the initial test.</p> <p>The positive control chemicals were tested concurrently. 2-Aminoanthracene was tested on all strains in the presence of rat and hamster S-9, 4-nitro-o-phenylenediamine was tested on TA 98 without S-9. Also without S-9, sodium azide was tested on TA100 and TA1535, and 9-aminoacridine was tested on TA1537.</p>
Results	Negative
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	Yes
Remarks for results	Limonene was inactive in <i>Salmonella</i> strains TA 1535, TA1537, TA98, and TA100 in the presence and absence of metabolic activation system.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.

References

Haworth, S., Lawlor T., Mortelmans K., Speck W., and Zeiger E. (1983) Salmonella Mutagenicity Test Results for 250 Chemicals. Environmental Mutagenesis Supplement 1, 3-142.

Substance Name	Orange peel oil, sweet (Citrus sinensis (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	Ames test
Test Type	Ames reverse mutation
System of Testing	Bacterial
GLP	No
Year	1989
Species/Strain	<i>Salmonella typhimurium</i> TA 98, TA 100, TA 1535, TA 1537, and TA1538
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	5000 micrograms/plate
Remarks for Test Conditions	The bacteria were cultured in Oxoid medium for approximately 12 hours. Selection of histidine revertants was performed in Vogel Bonner medium containing 2% glucose and 1.5% bacto agar. The preparation and composition of the overlay agar and a metabolic activation mix consisting of a 9000x g supernatant fraction (S9) of Aroclor 1254-induced adult male Sprague Dawley rat liver and necessary cofactors. The assays were conducted by addition of the test substance to a 2.0 ml volume of top agar along with a 0.1 ml aliquot of the bacterial culture and either the metabolic activation mix or an equivalent volume of phosphate buffer. These components were mixed rapidly and poured onto the minimal medium plates. After two days incubation at 37 °C, revertant colonies were counted. After two days incubation at 37 °C, revertant colonies were counted.
Results	No mutagenic effects.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Orange oil was inactive in <i>Salmonella</i> strains TA 1535, TA 1537, TA 1538, TA 98 & TA 100 both in the presence and absence of metabolic activation system.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Heck, J. D., Vollmuth, T. A., Cifone, M. A., Jagannath, D. R., Myhr B., and R.D. Curren (1989). An evaluation of food

flavoring ingredients in a genetic toxicity screening battery The Toxicologist, 9(1), 257.

Substance Name	Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	Ames test
Test Type	Ames reverse mutation
System of Testing	Bacterial
GLP	Ambiguous
Year	1990
Species/Strain	<i>Salmonella typhimurium</i> TA 98 and TA 100
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	100 microliters/plate
Remarks for Test Conditions	The Salmonella/microsome plate incorporation assay was performed. The test substances were tested in triplicates at several doses with a semilog interval up to a toxic or inhibitory dose or to a maximum of 100 ul/plate. Sterile distilled water was used as a solvent. Positive controls were 4-nitro-o-phenyldiamine (5 ug/plate for strains TA98 and TA1538) and mytomycin-C (1 ug/plate for strain TA94) S-9 level used was 50 microliters/plate.
Results	No mutagenic effects.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Orange oil was inactive in <i>Salmonella</i> strains TA 98 & TA 100 both in the presence and absence of metabolic activation system.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Crebelli R., Aquilina G., Conti L. and Carere A. (1990) Microbial mutagenicity screening of natural flavoring substances. Microbiologica 13, 115-119.
Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5

Method/guideline	Mouse Lymphoma Forward Mutation Assay (MLY)
Test Type	Forward mutation
System of Testing	Mammalian
GLP	Ambiguous
Year	1990
Species/Strain	L5178Y Mouse Lymphoma cell line
Metabolic Activation	With and without rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	100 micrograms/ml
Remarks for Test Conditions	The TK+/- -3.7.2C heterozygote of the L5178Y mouse lymphoma cell line was maintained in Fischer's medium containing 10% horse serum, antibiotics, glutamine, sodium pyruvate, and Pluronic F68. In a typical assay procedure, the thymidine kinase competent heterozygote was exposed to the test article in both the presence and absence of an induced rat liver S9 and cofactors. Cells were exposed for four hours, and then washed and incubated at 37 degrees for 48 hours before cloning. Colonies were counted after 10-14 days growth using an automatic colony counter. Mutant frequency was determined by calculating the ratio of mutant to viable colonies cloned without selective medium.
Results	No mutagenic effects.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Limonene was negative for mutagenic activity.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Heck J. D., Vollmuth, T. A., Cifone, M. A., Jagannath, D. R., Myhr B., and R.D. Curren (1989). An evaluation of food flavoring ingredients in a genetic toxicity screening battery The Toxicologist, 9(1), 257.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Mouse Lymphoma Forward Mutation Assay (MLY).
Test Type	Forward mutation

System of Testing	Mammalian
GLP	Ambiguous
Year	1989
Species/Strain	L5178Y Mouse Lymphoma cell line
Metabolic Activation	With and without rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	0-100 micrograms/ml
Remarks for Test Conditions	The highest dose of compound tested was determined by solubility or toxicity but did not exceed 5 mg/ml. All doses were tested in duplicate. Cells were exposed for four hours, and then washed and incubated at 37 °C for 48 hours before cloning. Colonies were counted after 9-12 days growth using an automatic colony counter.
Results	No mutagenic effects.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Limonene was negative for mutagenic activity.
Conclusion Remarks	No evidence of mutagenicity.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Myhr B., McGregor D., Bowers L., Riach C., Brown A.G., Edwards I., McBride D., Martin R., and Caspary W.J. (1990) L5178Y Mouse Lymphoma Cell Mutation Assay Results With 41 Compounds. Environmental and Molecular Mutagenesis. 16, 138-167.

Substance Name	Orange peel oil, sweet (Citrus sinensis (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	Mouse Lymphoma Forward Mutation Assay (MLY)
Test Type	Forward mutation
System of Testing	Mammalian
GLP	Ambiguous
Year	1990
Species/Strain	L5178Y Mouse Lymphoma cell line

Metabolic Activation	With and without rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	40-120 micrograms/ml, 125-200 micrograms/ml
Remarks for Test Conditions	The TK+/- -3.7.2C heterozygote of the L5178Y mouse lymphoma cell line was maintained in Fischer's medium containing 10% horse serum, antibiotics, glutamine, sodium pyruvate, and Pluronic F68. In a typical assay procedure, the thymidine kinase competent heterozygote was exposed to the test article in both the presence and absence of an induced rat liver S9 and cofactors. Cells were exposed for four hours, and then washed and incubated at 37 degrees for 48 hours before cloning. Colonies were counted after 10-14 days growth using an automatic colony counter. Mutant frequency was determined by calculating the ratio of mutant to viable colonies cloned without selective medium.
Results	Positive responses were observed with and without S-9, the latter only at highly toxic concentrations (relative survival less than 10%).
Cytotoxic concentration	Not given
Genotoxic Effects	Positive
Appropriate statistical evaluations	None given
Remarks for results	Orange oil was positive with and without S-9.
Conclusion Remarks	The authors stated that the low pH associated with the material may have contributed to the positive outcome. The authors recommended additional assays to clarify the positive response in the MLY assay.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Heck J. D., Vollmuth, T. A., Cifone, M. A., Jagannath, D. R., Myhr B., and R.D. Curren (1989). An evaluation of food flavoring ingredients in a genetic toxicity screening battery <i>The Toxicologist</i> , 9(1), 257.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Sister Chromatid Exchange in Chinese hamster ovary cells (Galloway <i>et al.</i> , 1985) with minor modifications
Test Type	Sister Chromatid Exchange
System of Testing	Mammalian
GLP	Ambiguous
Year	1990
Species/Strain	Chinese hamster ovary cells

Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats
Doses/Concentration	16.2-162 micrograms/ml limonene
Statistical Methods	Trend test (Margolin <i>et al.</i> , 1986)
Remarks for Test Conditions	The standard protocol as published by Galloway <i>et al.</i> was employed with minor modifications. Limonene was tested with and without activation. S-9 was used at a concentration of 15 or 20 microliters per milliliters.
Results	Negative
Cytotoxic concentration	Not given
Genotoxic Effects	Negative
Appropriate statistical evaluations	Yes
Remarks for results	Negative
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	Anderson B.E., Zeiger E., Shelby M.D., Resnick M.A., Gulati D.K., Ivett J.L., and Loveday K.S. (1990) Chromosome aberration and Sister Chromatid Exchange Test Results with 42 chemicals. Environmental and Molecular Mutagenesis 16, 55-137.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	In Vitro Sister Chromatid Exchange Test with Human Lymphocytes performed according to (Preston <i>et al.</i> , 1987)
Test Type	Sister Chromatid Exchange
System of Testing	Mammalian
GLP	Ambiguous
Year	1991
Species/Strain	Human lymphocytes
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats
Doses/Concentration	Up to 1000 micrograms/ml
Statistical Methods	Not given
Remarks for Test Conditions	Peripheral blood samples were obtained from one male and one female non-smoker. The test substances were added after 48 hr either for a period of 24 hours (without S-9 mix) or for a

Results	period of 2 hours (with S-9 mix). Myrcene was dissolved in ethanol. The mitotic index was determined for 1000 cells and given as number of mitoses per 1000 cells. No increase in frequency of sister chromatid exchange with or without metabolic activation.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Negative
Conclusion Remarks	The test substance was reported to have reduced the SCE inducing effect of S9 mix activated cyclophosphamide in human lymphocytes in a dose-dependent manner.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Kauderer B., Zamith H., Paumgarten J.R., and Speit G. (1991) Evaluation of the mutagenicity of <i>beta</i> -myrcene in mammalian cells in vitro. Environmental and Molecular Mutagenesis 18:28-34.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	In Vitro Sister Chromatid Exchange Test with Chinese Hamster Ovary V79 and HTC cells
Test Type	Sister Chromatid Exchange
System of Testing	Mammalian
GLP	Ambiguous
Year	1991
Species/Strain	Chinese hamster ovary cells
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats
Doses/Concentration	Up to 500 micrograms/ml
Statistical Methods	Not given
Remarks for Test Conditions	V79 cells were exposed to myrcene for three hours in the presence or absence of S-9 mix. Hepatic tumour cell (HTC) line cells were cultivated for the duration of one cell cycle in bromodeoxyuridine containing medium and for another 20 hours in bromodeoxyuridine free medium. HTC cells were treated with the test substances during the first cell cycle (20 h).
Results	Negative

Cytotoxic concentration	Not given
Genotoxic Effects	Negative
Appropriate statistical evaluations	Not given
Remarks for results	Negative
Conclusion Remarks	The test substance was reported to have reduced the SCE inducing effect of S9 mix activated cyclophosphamide and alfatoxin B1 in V79 and HTC cells in a dose-dependent manner.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Roscheisen C., Zamith H., Paumgarten F., and Speit G. (1991) Influence of <i>beta</i> -myrcene on sister chromatid exchanges induced by mutagens in V79 and HTC cells. Mutation Research 264, 43-49.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Chromosome Aberration in Chinese hamster ovary cells (Galloway <i>et al.</i> , 1985) with minor modifications
Test Type	Chromosomal Aberration assay
System of Testing	Mammalian
GLP	Ambiguous
Year	1990
Species/Strain	Chinese hamster ovary cells
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats
Doses/Concentration	50-500 micrograms/ml
Statistical Methods	Trend test (Margolin <i>et al.</i> , 1986)
Remarks for Test Conditions	The standard protocol as published by Galloway <i>et al.</i> was employed with minor modifications. Limonene was tested with and without activation. S-9 was used at a concentration of 15 or 20 microliters per milliliters. In earlier studies, 100 cells per dose were scored. In later studies, 200 cells per dose were scored. Fewer cells were scored if a strong response was observed or a dose was very toxic. Cells were scored for "simple (chromatid gaps and breaks, fragments, deletions, chromosome gaps and breaks, and double minutes), "complex" (interstitial deletions, triradials, quadriradials, rings, and dicentrics) and "other")pulverized, polyploids and endoreduplications) aberrations. The categories were combined to form a "total" category. Statistical analyses was conducted on the total category. The data were evaluated for both trend and dose point increase over the solvent control. A

binomial sampling assumption was used to evaluate an absolute increase in aberrations over the solvent control. Dose points with P values adjusted by Dunnett's method were considered to be significant if <0.05, but a trend was considered significant if P<0.003. If a trial had a positive trend and no significant doses, or if there was no trend and only one significant dose, the trial was judged equivocal (/); if a trial had significant trend and one significant dose it was judged positive (+W); and if the trial had two significant doses it was judged positive (+), whether or not a positive trend was obtained. Positive responses were generally repeated, whereas repeats were not required for the trials concluded to be negative. If positive responses were obtained both without and with S9, the laboratories were required to repeat only one activation condition.

Results	Negative
Cytotoxic concentration	Not given
Genotoxic Effects	Negative
Appropriate statistical evaluations	Yes
Remarks for results	Negative
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	Anderson B.E., Zeiger E., Shelby M.D., Resnick M.A., Gulati D.K., Ivett J.L., and Loveday K.S. (1990) Chromosome aberration and Sister Chromatid Exchange Test Results with 42 chemicals. Environmental and Molecular Mutagenesis 16, 55-137.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	<i>In Vitro</i> Chromosome Aberration Test with Human Lymphocytes performed according to (Preston <i>et al.</i> , 1987)
Test Type	Chromosomal Aberration assay
System of Testing	Mammalian
GLP	Ambiguous
Year	1991
Species/Strain	Human lymphocytes
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats
Doses/Concentration	Up to 1000 micrograms/ml

Statistical Methods	Not given
Remarks for Test Conditions	Peripheral blood samples were obtained from one male and one female non-smoker. The test substances were added after 48 hr either for a period of 24 hours (without S-9 mix) or for a period of 2 hours (with S-9 mix). Myrcene was dissolved in ethanol. The mitotic index was determined for 1000 cells and given as number of mitoses per 1000 cells.
Results	No induction of chromosome aberrations with or without metabolic activation. No indication of cytotoxicity.
Cytotoxic concentration	Not given
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Negative
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Kauderer B., Zamith H., Paumgarten J.R., and Speit G. (1991) Evaluation of the mutagenicity of <i>beta</i> -myrcene in mammalian cells in vitro. Environ. and Molecular Mutagenesis 18:28-34.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Cell Transformation in Syrian Hamster Embryo Cells
Test Type	Cell Transformation and Effect on Gap Junction Intercellular Communication
System of Testing	Mammalian
GLP	Ambiguous
Year	2000
Species/Strain	Syrian Hamster Embryo Cells
Metabolic Activation	None
Doses/Concentration	0.1-3 millimolar for Cell Transformation and 0.01-1 millimolar for Cellular Communication
Statistical Methods	One sided chi-square and ANOVA
Remarks for Test Conditions	Primary cell cultures from Syrian hamster embryos were prepared at 14 days gestation. The cells were exposed for 7 days starting 1 day after seeding of the target cells. Cells were observed for morphological transformation. Gap junction intercellular communication was measured as spreading of microinjected Lucifer Yellow dye to neighboring cells in a monolayer after 4 hours of exposure to test substances. The numbers of dye-coupled cells were counted 5-8 minutes following injection.

Results	Cell Transformation- 0.13% transformation frequency with a p-value of 0.089 using a chi-square.
Cytotoxic concentration	Cell Communication-No apparent effects Not given
Genotoxic Effects	Not applicable
Appropriate statistical evaluations	Yes
Remarks for results	Not statistically significant for cell transformation in Syrian Hamsters at the $\alpha = 0.05$ level. No effect on cellular communication.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Rivedal E., Mikalsen S.O., and Sanner T. (2000) Morphological transformation and effect on gap junction intercellular communication in Syrian Hamster Embryo Cells as Screening Tests for Carcinogens Devoid of Mutagenic Activity.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Cell Transformation in Syrian Hamster Embryo Cells
Test Type	Cell Transformation
System of Testing	Mammalian
GLP	No
Year	1980
Species/Strain	Syrian Hamster Embryo Cells
Metabolic Activation	None
Doses/Concentration	0.1-100 micrograms/ml
Statistical Methods	Not given
Results	Negative
Cytotoxic concentration	Not given
Genotoxic Effects	Negative
Appropriate statistical evaluations	None
Remarks for results	Negative
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.

References

Pienta R.J. (1980) Evaluation and relevance of the syrian hamster embryo cell system. The Predictive Value of Short Term Screening Tests in Carcinogenicity Evaluation, 149-160.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	V79-HPRT Gene Mutation Test with CHO cells according to Li <i>et al.</i> , 1987
Test Type	Gene mutation
System of Testing	Mammalian
GLP	Ambiguous
Year	1991
Species/Strain	Chinese hamster ovary cells
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced Sprague Dawley rats
Doses/Concentration	Up to 1000 micrograms/ml
Statistical Methods	Not given
Remarks for Test Conditions	The cells were exposed to myrcene or the control substances for 3 hours in the presence or absence of S-9 mix.
Results	No increase in mutation frequencies at the hprt-locus in V79 cells. No indication of induced cytotoxicity.
Cytotoxic concentration	Non-toxic up to 1000 microrams/ml
Genotoxic Effects	None
Appropriate statistical evaluations	None given
Remarks for results	Negative
Conclusion Remarks	The test substance was reported to have toxic and mutagenic effect of cyclophosphamide in V79 cells.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Kauderer B., Zamith H., Paumgarten J.R., and Speit G. (1991) Evaluation of the mutagenicity of <i>beta</i> -myrcene in mammalian cells in vitro. Environmental and Molecular Mutagenesis 18:28-34.

Substance Name	Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck)
CAS No.	8008-57-9

Method/guideline	<i>Bacillus subtilis</i> recessive assay
Test Type	Reverse mutation test
System of Testing	Bacterial
GLP	Ambiguous
Year	1989
Species/Strain	<i>Bacillus subtilis</i>
Metabolic Activation	Rat liver microsome fraction S9 from Aroclor induced rats
Doses/Concentration	5-30 microliters/plate
Results	No mutagenic effects.
Cytotoxic concentration	Not given
Genotoxic Effects	Negative
Appropriate statistical evaluations	None given
Remarks for results	Orange oil was negative with or without S-9 activation.
Data Qualities Reliabilities	Reliability code 3. Not reliable.
Remarks for Data Reliability	Code 3. Documentation insufficient for assessment.
References	Kuroda K., Yoo S., Ishibashi T. (1989) Rec-assay of natural food additives. Seikatsu Eisei 33, 15-23.

4.2.2 In vivo Genotoxicity

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Mammalian spot test
GLP	Ambiguous
Year	1984
Species/Strain	Mouse
Sex	Not reported
Route of Administration	Intraperitoneal
Doses/Concentration	215 mg/kg
Exposure Period	One day

Remarks for Test Conditions	Mouse embryos were treated in utero with limonene on days 10 and 11 post conception.
Appropriate statistical evaluations?	Yes, t-test
Genotoxic effects	No effects
Remarks for Results	No effects
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Fahrig R. (1984) Genetic mode of action of carcinogens and tumor promoters in yeast and mice. Molecular and General Genetics 194, 7-14.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	<i>in vivo</i> Cytogenetic Bone Marrow Assay
GLP	Ambiguous
Year	2001
Species/Strain	Mouse/B6C3F1
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/Concentration	0, 1000, or 2000mg/kg
Exposure Period	90 days
Remarks for Test Conditions	At the end of the 13-week subchronic study, a blood sample is obtained from male and female mice in each dose groups given either 1000 or 2000 mg/kg daily by oral gavage (usually 10 animals per treatment group per sex) and slides are prepared, fixed and stained as for the bone marrow studies. Sample collection time is typically between 0 and 24 hours. 1,000 to 10,000 mature erythrocytes (normochromatic erythrocytes or NCEs) are scored per animal for presence of micronuclei. These mature erythrocytes represent about 95% or more of the circulating erythrocytes. The percent PCE is determined in the blood as a measure of chemical-induced toxicity to the bone marrow. All data are analyzed separately for male and female mice. The acridine orange staining procedure that is used for micronucleus slides allows the scorer to differentiate between the recently formed, immature erythrocytes (polychromatic or PCE) that are less than 48 hr old, and mature erythrocytes 2-35 days old (normochromatic or NCE) based on their staining characteristics.

Appropriate statistical evaluations?
Effect on mitotic index or PCE/NCE ratio by dose level and sex

Yes, Kruskal-Wallis test, Mann-Whitney test

		Dose (g/kg)	No. of Animals	Mean MN-NCE/1000 NCE	Pairwise P
Vehicle Control:	Other Solvent	0.000	5	1.30 ± 0.34	
Test Chemical:		0.000	5	0.40 ± 0.19	0.5000
		1.000	10	1.35 ± 0.24	0.4555
		1.000	10	1.35 ± 0.24	0.4555
		2.000	2	1.25 ± 0.75	

Study ID and Trial No.	Start Date	Sample Time	Sex	Cell	Protocol	Trend Test P
Study ID A06528 Trial 2	11/11/2001	24 Hours	Male	NCE	GAV x 65, 90 Days	0.052

		Dose (g/kg)	No. of Animals	Mean MN-NCE/1000 NCE	Pairwise P
Vehicle Control:	Other Solvent	0.000	5	1.00 ± 0.22	
Test Chemical:		0.000	5	0.40 ± 0.19	0.5000
		1.000	10	1.20 ± 0.19	0.3137
		1.000	10	1.20 ± 0.19	0.3137
		2.000	1	3.00 ± 0.00	

Genotoxic effects

No statistically significant increases in frequency of micronuclei in either test groups of male or female mice

Remarks for Results

No dose-related increase in the mitotic index in bone marrow cells was reported for B6C3F1 mice administered 1000 or 2000 mg/kg of *beta*-myrcene daily by gavage for 90 days

Conclusion Remarks

The authors concluded that *beta*-myrcene was not clastogenic to mice when orally administered at dose levels up to 2000 mg/kg bw.

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability References

Code 1. Published in peer reviewed journal and standard protocols were used.
National Toxicology Program (NTP Draft) (2003) Initial study results from a 90 day toxicity study on *beta*-myrcene in mice and rats. Study number C99023 and A06528.

Substance Name

Myrcene

CAS No.

123-35-3

Remarks for Substance

beta-Myrcene

Method/guideline	in vivo Cytogenetic Bone Marrow Assay																																												
GLP	Ambiguous																																												
Year	1993																																												
Species/Strain	Rat/Wistar																																												
Sex	Male and Female																																												
Route of Administration	Oral-Gavage																																												
Doses/Concentration	100, 500 or 1000 mg/kg																																												
Exposure Period	24 and 48 hours																																												
Remarks for Test Conditions	beta-Myrcene (100, 500 or 1000 mg/kg) was orally administered via gavage to two or four male and female Wistar rats. Corn oil was used as the negative control while cyclophosphamide (30 mg/kg via intraperitoneal injection) was used as the positive control. A mitotic inhibitor (colchicine 5 mg/kg ip) was injected 1 hr before sacrifice. At 24 or 48 hours, animals were sacrificed and bone marrow cells harvested. Evaluations included the mitotic index and the frequency of chromosomal aberrations.																																												
Appropriate statistical evaluations?	Yes, Kruskal-Wallis test, Mann-Whitney test																																												
Effect on mitotic index or PCE/NCE ratio by dose level and sex	<p>Corn oil (24 hr sampling)</p> <table border="0"> <thead> <tr> <th>Number</th> <th>Sex</th> <th>Mitotic index</th> </tr> </thead> <tbody> <tr> <td>2</td> <td>Males</td> <td>11.5</td> </tr> <tr> <td>2</td> <td>Females</td> <td>12.5</td> </tr> <tr> <td>4</td> <td>Males and Females</td> <td>12.0 +/- 2.9</td> </tr> </tbody> </table> <p>beta-Myrcene</p> <table border="0"> <tbody> <tr> <td>100 mg/kg bw (24 hr)</td> <td></td> </tr> <tr> <td>2 Males</td> <td>14.0</td> </tr> <tr> <td>2 Females</td> <td>17.0</td> </tr> <tr> <td>4 Males and Females</td> <td>15.5 +/- 8.5</td> </tr> </tbody> </table> <p>500 mg/kg bw (24 hr)</p> <table border="0"> <tbody> <tr> <td>2 Males</td> <td>19.5</td> </tr> <tr> <td>2 Females</td> <td>19</td> </tr> <tr> <td>4 Males and Females</td> <td>19.2 +/-1.7</td> </tr> </tbody> </table> <p>1000 mg/kg (24 hr)</p> <table border="0"> <tbody> <tr> <td>4 Males</td> <td>21.0</td> </tr> <tr> <td>4 Females</td> <td>23.0</td> </tr> <tr> <td>4 Males and Females</td> <td>22.0 +/- 6.1</td> </tr> </tbody> </table> <p>1000 mg/kg bw (48 hr)</p> <table border="0"> <tbody> <tr> <td>2 Males</td> <td>16.5</td> </tr> <tr> <td>2 Females</td> <td>12.0</td> </tr> <tr> <td>4 Males and Females</td> <td>14.2 +/-3.6</td> </tr> </tbody> </table> <p>Cyclophosphamide (24 hr)</p> <table border="0"> <tbody> <tr> <td>30 mg/kg ip</td> <td></td> </tr> <tr> <td>2 Males</td> <td>10.5</td> </tr> <tr> <td>2 Females</td> <td>8.5</td> </tr> </tbody> </table>	Number	Sex	Mitotic index	2	Males	11.5	2	Females	12.5	4	Males and Females	12.0 +/- 2.9	100 mg/kg bw (24 hr)		2 Males	14.0	2 Females	17.0	4 Males and Females	15.5 +/- 8.5	2 Males	19.5	2 Females	19	4 Males and Females	19.2 +/-1.7	4 Males	21.0	4 Females	23.0	4 Males and Females	22.0 +/- 6.1	2 Males	16.5	2 Females	12.0	4 Males and Females	14.2 +/-3.6	30 mg/kg ip		2 Males	10.5	2 Females	8.5
Number	Sex	Mitotic index																																											
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4 Males and Females 9.5+/- 3.7

Genotoxic effects	2 Females 4 Males and Females Dose related increase in mitotic index, no statistically significant increases in frequency of chromosomal aberrations
Remarks for Results	A dose-related increase in the mitotic index in bone marrow cells was reported for rats administered beta-myrcene. The authors commented that this be an interaction between beta-myrcene, which is known to induce CYP-P450 enzymes, and colchicine, which arrests cell division at metaphase. Myrcene may have increased the bioavailability of colchicine leading to the increase in mitotic index observed in the experiment. No significant increases in chromosomal aberrations were reported in the treated animals at either 24 or 48 hours.
Conclusion Remarks	The authors concluded that given the results, beta-myrcene was not clastogenic to the rat when orally administered at dose levels up to 1000 mg/kg bw.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability References	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles. Zamith H.P., Vidal M.N.P., Speit G. and Paumgarten F.J.R. (1993) Absence of genotoxic activity of beta-myrcene in the in vivo cytogenetic bone marrow assay. Brazilian J Med Biol Res 26, 93-98.

4.3 Repeat dose Toxicity

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	beta-Myrcene; greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study
GLP	Yes
Year	1990
Species/strain	F344/N Rats
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 250, 500, 1000, 2000, 4000 mg/kg bw/d
Exposure Period	13 weeks; 3 weeks/2 days (Special Study)
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes

Post Exposure

Remarks for Test Conditions	<p>Groups of ten rats of each sex were administered 0, 250, 500, 1000, 2000, 4000 mg/kg bw/d mg/kg beta-myrcene in corn oil by gavage once per day, five days a week for 13 weeks. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week. Necropsies were performed on all animals. Histological examinations were performed on all vehicle control and high dose animals and all female rats in the 1200 mg/kg group. Tissues examined included adrenal glands, brain, colon, esophagus, eyes (if grossly abnormal), femur, sternbrae or vertebrae including marrow, gross lesions and tissue masses with regional lymph nodes, heart kidneys, liver, lungs and mainstem bronchi, mammary gland, mandibular or mesenteric lymph nodes, pancreas, parathyroids, pituitary gland, prostate/testes or ovaries/uterus, salivary glands, small intestine, spinal cord (if neurologic signs present), spleen, stomach, thymus, thyroid gland, trachea, and urinary bladder. Kidneys were examined in all male rats.</p> <p>Right kidneys of male rats were frozen while left kidneys were processed for Mallory Heidenhain staining and H&E staining for investigation of alpha-2-globulin in male rats. Also, a Special Study Groups (10/dose/sex) was given three doses of β-myrcene daily for 3 weeks and 2 days. Body weights were measured weekly and hematological examinations and blood chemical determination were performed at termination on Day 23. At termination, the left kidneys were frozen and the right kidneys were processed and microscopically examined for the presence of hyaline droplets using Mallory Heidenhain staining and H&E staining.</p>
NOAEL (NOEL)	Not assigned
LOAEL(LOEL)	250 mg/kg bw/d
Toxic Response/effects by Dose Level	<p>All animals in the Core group and Special Study groups at 4000 mg/kg bw per day group died within the first 12 days of the study. At 2000 mg/kg bw per day, similar survival was reported in both the Core (2/10) and Special Groups (4/10) of males and females (4/10 in each group). In animals that died prior to study termination, clinical signs included lethargy, ruffled fur, abnormal breathing, or thin appearance. Greater than 10% decrease in mean body weight gain was recorded for males at the 1000, 2000, and 4000 mg/kg bw per day dose levels. Group mean body weights for females in all dose groups and in Special Study groups were within 10% of those reported for respective control groups.</p> <p>Hematological examination revealed decreases in white blood cells (27% and 24%) and lymphocytes (35% and 25%) in males at Day 93 and females at Day 23 given 2000 mg/kg bw per day. Increases in reticulocytes were reported in males at the 1000 and 2000 mg/kg bw per day dose levels. A dose-dependent increase in absolute and relative liver and kidney weights was reported in males and females rats compared to that for</p>

respective control groups. A dose-dependent decrease in mean thymus weight beginning at 500 mg/kg bw per day was reported in males. A decrease in mean thymus weight was also reported in females but only in the highest dose group. Clinical chemistry determinations of the kidney (BUN and creatinine) and liver (ALT and SDH) parameters showed no significant difference between test and control groups.

In males, histopathological examination revealed consistent evidence of renal tubular hyaline droplet formation in dose groups surviving to Day 93 (250, 500, and 1000 mg/kg bw per day) and in control males (10/10). Renal tubular loading of droplets was scored on a subjective scale of 0 to 4.0. Both control and the three low dose groups gave a score of 3.0 after 93 days. Control and test animals all showed evidence of nephropathy with increasing severity reported in a dose related manner. Control and test groups showed evidence of porphyrin pigmentation of the Harderian gland. The incidence of this latter effect also increased with dose. There was no consistent evidence of histopathology of the liver in the 250, 500, or 1000 mg/kg bw per day dose groups. Based on the alpha_{2u}-globulin renal tubular effects observed at all dose levels and in the control group, a no effect level could not be assigned.

In females, control and test animals all showed evidence of nephropathy with increasing incidence and severity reported in a dose related manner. Control (3/10) and test groups 4/10, 2/10, and 4/10 at 250, 500, and 1000 mg/kg bw per day, respectively) showed evidence of porphyrin pigmentation of the Harderian gland. The incidence of this latter effect also increased with dose. All test groups showed evidence of renal tubular degeneration while dose levels of 1000 mg/kg bw per day and above exhibited splenic atrophy, olfactory epithelial degeneration and chronic nasal irritation. Based on the presence of renal tubular degeneration, a no effect level could not be assigned.

Statistical Evaluation

Yes, Kaplan-Meier used for probability of survival. Statistical analyses used for possible dose-related effect on survival was Cox (1972) for testing two groups for equality; and Tarone's (1975) life table test for a dose-related trend.

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Guideline study.

References

National Toxicology Program (NTP Draft) (2003) Initial study results from a 90 day toxicity study on beta-myrcene in mice and rats. Study number C99023 and A06528.
Tarone R.E. (1975) Tests for trend in life table analysis. *Biometrika* 62; 679-682.
Cox D.R. (1972) Regression models and life tables. *J.R. Stat. Soc. B34*: 187-220.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	beta-Myrcene ; greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study
GLP	Yes
Year	1990
Species/strain	B6C3F1 Mice
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 250, 500, 1000, 2000, 4000 mg/kg bw/d
Exposure Period	13 weeks;
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Post Exposure	
Remarks for Test Conditions	Groups of ten mice of each sex were administered 0, 250, 500, 1000, 2000, 4000 mg/kg bw/d mg/kg beta-myrcene in corn oil. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week. Necropsies were performed on all animals. Histological examinations were performed on all vehicle control and high dose animals. Tissues examined included adrenal glands, brain, colon, esophagus, eyes (if grossly abnormal), femur or sternbrae or vertebrae including marrow, gallbladder, gross lesions and tissue masses with regional lymph nodes, heart kidneys, liver, lungs and mainstem bronchi, mammary gland, mandibular or mesenteric lymph nodes, pancreas, parathyroids, pituitary gland, prostate/testes or ovaries/uterus, salivary glands, small intestine, spinal cord (if neurological signs present), spleen, stomach, thymus, thyroid gland, trachea, and urinary bladder.
NOAEL (NOEL)	Based on these data a NOEL for exposure to β -myrcene in B6C3F1 female mice for 14 weeks is 250 mg/kg bw per day and an NOAEL for male B6C3F1 mice is 500 mg/kg bw per day. Females: 250 mg/kg bw/d Males: 500 mg/kg bw/d
LOAEL(LOEL)	Females: 500 mg/kg bw/d Males: 1000 mg/kg bw/d
Toxic Response/effects by Dose Level	All animals in the 4000 mg/kg bw per day group died within the first three days while 9 of 10 males and 8 of ten females died prior to Week 5. In animals that died prior to study termination,

clinical signs included lethargy, abnormal breathing, or thin appearance. Group mean body weight gains were depressed for both males (-22.5%) and females (-2.4%) in the 1000 mg/kg bw per day group.

Mean female body weights of the 250 and 500 mg/kg bw per day dose levels were either unchanged or slightly depressed while the 1000 and 2000 mg/kg bw per day was significantly less than that of the respective control groups. For males, relative liver weights were increased for the 1000 and 2000 mg/kg bw per day groups only. For females, absolute and relative liver weights increased in a dose related manner at dose levels of 500 mg/kg bw per day and above. Increased relative liver weights in females were partly due to lower body weight gain at the two lower dose levels. Hematological examination revealed small decreases (<6%) in red blood cells, hemoglobin, and hematocrit and increases (<3%) in mean corpuscular volume and mean corpuscular hemoglobin at the 1000 mg/kg bw per day level in both sexes. At the 2000 mg/kg bw per day level, these changes were more pronounced (5-43%). No significant differences in clinical chemistry parameters were reported between test and control animals. In females, centrilobular hypertrophy and necrosis of the liver and forestomach irritation was reported at the 2000 and 4000 mg/kg bw per day dose levels but not at lower dose levels. In males, minimal or mild centrilobular hypertrophy was reported in all dose groups with a single incidence in the 250 mg/kg bw per day group increasing to 100% in the 1000 mg/kg bw per day group. Liver necrosis was reported in the 1000 and 2000 mg/kg bw per day groups. Also in males, the 100% incidence of renal cytoplasmic vacuolation in the control group decreased to 10% in the 500 mg/kg bw per day group. Based on these data a NOEL for exposure to β -myrcene in B6C3F1 female mice for 14 weeks is 250 mg/kg bw per day and an NOAEL for male B6C3F1 mice is 500 mg/kg bw per day.

Statistical Evaluation

Yes, Kaplan-Meier used for probability of survival. Statistical analyses used for possible dose-related effect on survival was Cox (1972) for testing two groups for equality; and Tarone's (1975) life table test for a dose-related trend.

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Guideline study.

References

National Toxicology Program (NTP Draft) (2003) Initial study results from a 90 day toxicity study on beta-myrcene in mice and rats. Study number C99023 and A06528.

Tarone R.E. (1975) Tests for trend in life table analysis. *Biometrika* 62; 679-682.

Cox D.R. (1972) Regression models and life tables. *J.R. Stat. Soc. B34*: 187-220.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study NTP TR 347
GLP	Yes
Year	1990
Species/strain	B6C3F1 Mice
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 413, 825, 1650, 3300, or 6600 mg/kg bw/d
Exposure Period	16 days
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Post Exposure	4 days
Remarks for Test Conditions	Groups of five mice of each sex were administered 0, 413, 825, 1650, 3300, or 6600 mg/kg <i>d</i> -limonene in corn oil by gavage once per day for 12 days over a 16 day period. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week. Necropsies were performed on all animals.
NOAEL (NOEL)	1650 mg/kg bw/d
LOAEL(LOEL)	3300 mg/kg bw/d
Toxic Response/effects by Dose Level	See remarks for results.
Statistical Evaluation	Yes, Kaplan-Meier used for probability of survival. Statistical analyses used for possible dose-related effect on survival was Cox (1972) for testing two groups for equality; and Tarone's (1975) life table test for a dose-related trend.
Remarks for Results	All but one animal receiving 3300 or 6600 mg/kg bw/d limonene died within three days of study initiation. No treatment-related clinical signs were observed in mice receiving doses of 1650 mg/kg bw/d or lower.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	National Toxicology Program (NTP) (1990) Carcinogenicity and toxicology studies of <i>d</i> -limonene in F344/N Rats and B6C3F1 mice. NTP-TR-347. U.S. Dept of Health and Human Services. NIH Publication No. 90-2802.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study NTP TR 347
GLP	Yes
Year	1990
Species/strain	F344/N Rats
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 413, 825, 1650, 3300, or 6600 mg/kg bw/d
Exposure Period	16 days
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Post Exposure	4 days
Remarks for Test Conditions	Groups of five rats of each sex were administered 0, 413, 825, 1650, 3300, or 6600 mg/kg <i>d</i> -limonene in corn oil by gavage once per day for 12 days over a 16 day period. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week. Necropsies were performed on all animals.
NOAEL (NOEL)	1650 mg/kg bw/d
LOAEL(LOEL)	3300 mg/kg bw/d
Toxic Response/effects by Dose Level	All but two females receiving 3300 or 6600 mg/kg bw/d limonene died within two days of study initiation. No treatment related clinical signs were observed in rats receiving doses of 1650 mg/kg bw/d or lower.
Statistical Evaluation	Yes, Kaplan-Meier used for probability of survival. Statistical analyses used for possible dose-related effect on survival was Cox (1972) for testing two groups for equality; and Tarone's (1975) life table test for a dose-related trend.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	National Toxicology Program (NTP) (1990) Carcinogenicity and toxicology studies of <i>d</i> -limonene in F344/N Rats and B6C3F1 mice. NTP-TR-347. U.S. Dept of Health and Human Services. NIH Publication No. 90-2802.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study NTP TR 347
GLP	Yes
Year	1990
Species/strain	F344/N Rats
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 150, 300, 600, 1200, or 2400 mg/kg bw/d
Exposure Period	13 weeks
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Remarks for Test Conditions	Groups of ten rats of each sex were administered 0, 150, 300, 600, 1200 or 2400 mg/kg bw/d <i>d</i> -limonene in corn oil by gavage once per day, five days a week for 13 weeks. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week. Necropsies were performed on all animals. Histological examinations were performed on all vehicle control and high dose animals and all female rats in the 1200 mg/kg group. Tissues examined included adrenal glands, brain, colon, esophagus, eyes (if grossly abnormal), femur, sternbrae or vertebrae including marrow, gross lesions and tissue masses with regional lymph nodes, heart kidneys, liver, lungs and mainstem bronchi, mammary gland, mandibular or mesenteric lymph nodes, pancreas, parathyroids, pituitary gland, prostate/testes or ovaries/uterus, salivary glands, small intestine, spinal cord (if neurologic signs present), spleen, stomach, thymus, thyroid gland, trachea, and urinary bladder. Kidneys were examined in all male rats.
NOAEL (NOEL)	300 mg/kg bw/d
LOAEL(LOEL)	600 mg/kg bw/d
Toxic Response/effects by Dose Level	See remarks for results.
Statistical Evaluation	Yes, Kaplan-Meier used for probability of survival. Statistical analyses used for possible dose-related effect on survival was Cox (1972) for testing two groups for equality; and Tarone's (1975) life table test for a dose-related trend.

Remarks for Results

Ninety percent of female rats (9/10) and fifty percent of male rats (5/10) receiving 2400 mg/kg bw/d limonene died within the first week of the study. The final mean body weights of male rats receiving the three highest doses (600, 1200 or 2400 mg/kg bw/d) were reported to be 6%, 12%, or 23% lower than that of the controls, respectively. Rough hair coats, lethargy, and excessive lacrimation were observed for all animals at the two highest dose levels. Nephropathy was reported for all groups of male rats but a dose related increase in severity of the lesion was reported for the dosed groups. The nephropathy was characterized by degeneration of epithelium in the convoluted tubules, granular casts with tubular lumens, primarily in the outer stripe of the outer medulla, and regeneration of the tubular epithelium. Hyaline droplets were observed in the epithelium of the proximal convoluted tubules in all groups of male rats including vehicle controls. Upon further review to determine if there were differences in these findings between control and treated animals, the blinded slides revealed no definite differences in the accumulation of hyaline droplets.

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Guideline study.

References

National Toxicology Program (NTP) (1990) Carcinogenicity and toxicology studies of *d*-limonene in F344/N Rats and B6C3F1 mice. NTP-TR-347. U.S. Dept of Health and Human Services. NIH Publication No. 90-2802.
 Tarone R.E. (1975) Tests for trend in life table analysis. Biometrika 62; 679-682.
 Cox D.R. (1972) Regression models and life tables. J.R. Stat. Soc. B34: 187-220.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study NTP TR 347
GLP	Yes
Year	1990
Species/strain	B6C3F1 Mice
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 125, 250, 500, 1000 or 2000 mg/kg bw/d
Exposure Period	13 weeks

Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Remarks for Test Conditions	Groups of ten mice of each sex were administered 0, 125, 250, 500, 1000 or 2000 mg/kg bw/d <i>d</i> -limonene in corn oil by gavage once per day, five days a week for 13 weeks. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week. Necropsies were performed on all animals. Histological examinations were performed on all vehicle control and high dose animals. Tissues examined included adrenal glands, brain, colon, esophagus, eyes (if grossly abnormal), femur or sternbrae or vertebrae including marrow, gallbladder, gross lesions and tissue masses with regional lymph nodes, heart kidneys, liver, lungs and mainstem bronchi, mammary gland, mandibular or mesenteric lymph nodes, pancreas, parathyroids, pituitary gland, prostate/testes or ovaries/uterus, salivary glands, small intestine, spinal cord (if neurological signs present), spleen, stomach, thymus, thyroid gland, trachea, and urinary bladder.
NOAEL (NOEL)	500 mg/kg bw/d
LOAEL(LOEL)	1000 mg/kg bw/d
Toxic Response/effects by Dose Level	One of 10 males and 2/10 females administered 2000 mg/kg bw/d limonene and 1/10 females administered 500 mg/kg bw/d limonene died before the end of the study. Several other animals also died as a result of gavage error. Mean body weights were 10% lower than control for male mice and 2% lower than control for female mice for the two highest dose levels. An alveolar cell adenoma was reported in the lung of one female at the highest dose level. Clinical signs of rough hair coats and decreased activity were reported for the two highest dose levels.
Statistical Evaluation	Yes
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	National Toxicology Program (NTP) (1990) Carcinogenicity and toxicology studies of <i>d</i> -limonene in F344/N Rats and B6C3F1 mice. NTP-TR-347. U.S. Dept of Health and Human Services. NIH Publication No. 90-2802.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	<i>d</i> -limonene; greater than 99% pure
Method/guideline	Subchronic nephrotoxicity study
GLP	Ambiguous

Year	1989
Species/strain	F344/N Rats
Sex	Male
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 2, 5, 10, 30, or 75 mg/kg bw/d
Exposure Period	13 weeks
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Remarks for Test Conditions	Groups of five week old male rats received 0, 2, 5, 10, 30 or 75 mg/kg bw/d <i>d</i> -limonene daily via oral gavage for 13 weeks (5 days a week). Rats from selected dose groups were necropsied throughout the study (days 8-29), with all remaining rats necropsied at the end of the study. Rats were observed daily for toxicity signs. Body weights were taken daily.
NOAEL (NOEL)	5 mg/kg bw/d
LOAEL(LOEL)	30 mg/kg bw/d
Toxic Response/effects by Dose Level	5 mg/kg bw/d no effects 10 mg/kg bw/d formation of hyaline droplets 30 mg/kg bw/d increased relative kidney weight, formation of hyaline droplets; 75 mg/kg bw/d increased relative kidney and liver weights; increased granular casts in outer medulla; formation of hyaline droplets.
Statistical Evaluation	Yes, linear regression analyses
Remarks for Results	Linear regression analyses indicated increased relative kidney and liver weights at the two highest dose levels. Histological examination revealed changes characterized by hyaline droplet formation, granular casts and multiple cortical changes, all of which was classified as chronic nephrosis. Exacerbation of hyaline droplet formation was reported at the earliest necropsy eight days after administration at the 10 mg/kg bw/d dose level.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Webb D.R., Ridder M., and Alden C.L. (1989) Acute and subchronic nephrotoxicity of <i>d</i> -limonene in Fischer 344 rats.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
GLP	No
Year	1977

Species/strain	Rat/Sprague Dawley
Sex	Not reported
Route of Administration	Oral
Doses/concentration Levels	0, 277, 554, 1385, or 2770 mg/kg bw/d
Exposure Period	30 days
Frequency of Treatment	Daily
Control Group	Yes
Remarks for Test Conditions	The test substance was orally administered to Sprague-Dawley rats daily for 30 days at the following dose levels 0, 277, 554, 1385, or 2770 mg/kg bw in order to investigate the effect on the fine structure of the liver, kidney and blood cells.
NOAEL (NOEL)	1385 mg/kg bw/d
LOAEL(LOEL)	2770 mg/kg bw/d
Toxic Response/effects by Dose Level	No morphological changes of renal corpuscles and tubular cells were observed. Some alterations were detected in the glomerular epithelium from the kidneys of rats treated at the highest dose level.
Statistical Evaluation	Not given
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Kodama R., Tahara S., Sato K., Noda K., Ide H., Nishihara H. (1977b) Studies on <i>d</i> -limonene as a Gallstone Solubilizer Fine Structure of Liver, Kidneys and Blood Cells from Rats given <i>d</i> -limonene. Oyo Yakuri 13(6), 875.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study NTP TR 347
GLP	Yes
Year	1990
Species/strain	F344/N Rats
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	Males: 0, 75 or 150 mg/kg bw/d; Females: 0, 300 or 600 mg/kg bw/d

Exposure Period	103 weeks
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Remarks for Test Conditions	Groups of fifty male and fifty female rats each were administered 0, 75 or 150 mg/kg bw/d or 0, 300 or 600 mg/kg bw/d <i>d</i> -limonene, respectively, in corn oil by gavage once per day, five days a week for 103 weeks. Animals were housed five per cage and fed ad libitum. The animals were observed twice per day and weighed once per week for 12 weeks and once per month thereafter. Necropsies were performed on all animals. Histological examinations were performed on all animals dying during the study; all vehicle control; all low dose female rats and all high dose animals. Tissues examined included adrenal glands, brain, cecum, colon, costochondral junction, duodenum, epididymus/seminal vesicles/tunica vaginalis/scrotal sac/prostate/testes or ovaries/uterus, esophagus, eyes, femur or sternbrae or vertebrae including marrow, gross lesions and tissue masses with regional lymph nodes, heart, ileum, jejunum, kidneys, larynx and pharynx, liver, lungs and bronchi, mammary gland, mandibular or mesenteric lymph nodes, nasal cavity and turbinates, oral cavity, pancreas, parathyroids, pituitary gland, preputial or clitoral gland, rectum, salivary glands, sciatic nerve, skin, spinal cord, spleen, stomach, thigh muscle, thymus, thyroid gland, trachea, urinary bladder and Zymbal gland. Tissues examined in low dose male rat groups included adrenal glands, kidney, liver, spleen, and testis.
NOAEL (NOEL)	Undetermined (male); 300 mg/kg bw/d (female)
LOAEL(LOEL)	75 mg/kg bw/d (male); 600 mg/kg bw/d (female)
Toxic Response/effects by Dose Level	<p>Male Rats Body weight-5% reduction in high dose group. Survival rate-29/50; 33/50;40/50. Non-neoplastic effects-mineralization 7/50; 43/50; 48/50; epithelial hyperplasia 0/50; 35/50; 43/50 of the renal papilla; renal tubular cell hyperplasia 0/50; 4/50; 7/50. Neoplastic effects-renal tubular cell adenomas 0/50; 4/50; 8/50 an adenocarcinomas 0/50; 4/50; 3/50.</p> <p>Female Rats Body weight-5% reduction in high dose group. Survival rate-42/50; 40/50;26/50. No neoplastic or nonneoplastic effects.</p>
Statistical Evaluation	Yes
Remarks for results	Mean body weights for male rats administered 150 mg/kg bw/d <i>d</i> -limonene were generally 4-7% lower than vehicle controls from week 2 to study termination. Mean body weights of high dose females were generally 4-7% lower than vehicle controls from week 28 to study termination. No treatment related clinical signs were reported for the duration of the study. Survival of the high dose male group was significantly greater than that of the vehicle alone after week 81. Survival of the high dose female

group was significantly lower than that of the vehicle controls after week 39. In the kidneys of male rats, dose-related increases in the incidences of mineralization and epithelial hyperplasia. A dose-related increase in the severity of spontaneous nephropathy was reported in male rats administered limonene. Increased incidence of tubular cell hyperplasia and neoplasia was also reported in dosed male rats. Tubular cell adenoma incidence in high dose male rats and tubular cell adenoma or tubular cell carcinomas (combined) in dosed male rats were significantly greater than vehicle controls.

Conclusion Remarks

The authors determined that under the conditions of these 2-year gavage studies there was clear evidence of carcinogenic activity of *d*-limonene for male F344/N rats as shown by increased incidences in tubular cell hyperplasia, adenomas, and adeno-carcinomas of the kidney. There was no evidence of carcinogenic activity of *d*-limonene for female rats receiving 300 or 600 mg/kg bw/d.

It has been demonstrated that renal lesions, which were observed in the NTP study, resulted from the accumulation of aggregates of alpha-2u-globulin (a low molecular-weight protein synthesized in the liver) and limonene or its metabolites in the P2 segment of the renal proximal tubule. This phenomenon has only been observed in the male F344/N rat [Strasser, 1988; Borghoff *et al.*, 1990]. The gene that encodes alpha-2u-globulin has been isolated and the sequence deduced [Untermann *et al.*, 1981]. These proteins are expressed in the liver under hormonal control [Roy and Neuhaus, 1967; Wang and Hodgetts, 1998]. alpha-2u-Globulin belongs to the a-2m-globulin super family of proteins that are characterized by a unique hydrophobic binding pocket. The lesions do not develop in the female F344/N rat or in humans [Bucher *et al.*, 1986]. Subsequent investigations have shown that the alpha-2u-globulin nephropathy found in the F344/N male rat does not develop in mammals that do not express the hepatic form of a-2m-globulin [Swenberg, 1989] such as other strains of rats [Dietrich and Swenberg, 1991], mice [Bucher *et al.*, 1986; Lehman-McKeeman, 1994] and dogs [Webb, 1990].

Transgenic mice that express rat alpha-2u-globulin were tested for their ability to form hyaline droplets and develop nephropathies similar to their adult male rat counterparts [Lehman-McKeeman and Caudill, 1994]. This study involved male F344 rats as positive control, transgenic C57BL/6J mice as experimental group and native C57BL/6 mice as negative controls. The animals at age 70-75 days were placed in metabolic cages and received 150 mg/kg bw per day *d*-limonene in corn oil by gavage for three days. Limonene was used to induce renal nephropathy in adult male rats, as it was shown to be a potent inducer in the NTP studies [EPA, 1991; NTP, 1990]. Twenty-four (24) hours after the last dose, the animals were sacrificed and the kidneys analyzed for evidence of nephropathy. Hyaline droplet formation was evaluated on a subjective scale, size and intensity (0-4) multiplied by tubular loading (0-3) for an overall scale of 0-12 with 12 being the most

severe. In the absence of *d*-limonene, the control groups transgenic mice and rats showed a hyaline droplet score of 1+/- 0 and 6 +/- 0.5, respectively. The test transgenic mice and rats showed a hyaline droplet score of 2.5 +/- 0.3 and 11 +/- 1.3, respectively upon dosing with *d*-limonene. The native mice developed no signs of hyaline droplet formation and tested negative for presence of alpha-2u-globulin in their urine. The authors assert that based on the data presented "alpha-2u-globulin is the only protein that is involved in the etiology of hyaline droplet nephropathy".

An increase in the kidney-type- alpha-2u-globulin was seen in the urine of male Sprague-Dawley rats when these animals were administered greater than 30 mg/kg/day of *d*-limonene for 7 days by gavage. The increases in the urinary kidney-type-alpha-2u-globulin are dose-dependent and parallel-elevated accumulation in the kidney cells [Saito, 1996]

In another study, adult male Wistar rats were administered two groups of chemical compounds, including 138 mg/kg bw isophorone, potassium bromate, 2-propanol and a series of benzene and anthracene derivatives, to study induction of accumulation of alpha-2u-globulin and structure-activity relationships. A monoclonal antibody against alpha-2u-globulin was employed in a competitive ELISA procedure to determine its concentration in urine or tissue samples without purification. Plasma concentrations of alpha-2u-globulin were not significantly increased by any of the test compounds at 1 mmol/kg bw. Kidney tissue concentrations were found to be 297-300% higher than controls. The hyaline droplet accumulating (HDA) potential was dependent on the test compound but there was no relationship between HAD activity and the structure or the pathway used to metabolize the test substance [Hildebrand, 1997]

While humans produce low molecular weight serum proteins, which are reabsorbed by the kidney, there is no evidence that alpha-2u-globulin is produced [Olson, 1990]. Urine collected from adult male F344 rats and humans revealed no evidence indicative of alpha-2u-globulin production in humans [Olson, 1990].

It is unknown whether any human serum proteins possess a binding site similar to that of alpha-2u-globulin. Although this is a possibility, it appears remote, since female rats and mice do not show the renal changes noted in male rats exposed to limonene. It should be noted that there is a class of human proteins referred to as the alpha-2u-globulin -related proteins. They appear to have no functional relationship to the adult male rat urine proteins. The human protein has a higher molecular weight, 25 kDa and is a component of a neutrophil gelatinase complex [Kjeldsen *et al.*, 2000 and Triebel *et al.*, 1992]. An extensive review of the current scientific literature and genome databases reveals no native protein or biological entity that acts as a nephropathy agent like mature male rat alpha-2u-globulin. The accumulated evidence indicates that it is the unique

anatomical, physiological, and biochemical properties of the male rat kidney, especially the proximal convoluted tubule, that allows *d*-limonene to interfere with renal processing of the strain-specific alpha-2u-globulin. Therefore, this process is not predictive of human carcinogenicity. In a comprehensive review of alpha-2u-globulin nephropathy and associated renal tubule tumors produced in the male F344/N rat exposed to limonene and other simple chemical substances (e.g. isophorone, decalin and methyl isobutyl ketone), it was concluded that the F344/N rat is not an appropriate model for assessing human renal carcinogenic risk [EPA, 1991]. After careful review, it has been concluded that the mechanisms leading to the renal carcinogenic findings in the F344/N male rat are largely known and strongly indicate that the nephropathy associated with *d*-limonene have no significance for human risk assessment [Burdock *et al.*, 1990].

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Guideline study.

References

National Toxicology Program (NTP) (1990) Carcinogenicity and toxicology studies of *d*-limonene in F344/N Rats and B6C3F1 mice. NTP-TR-347. U.S. Dept of Health and Human Services. NIH Publication No. 90-2802.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	Greater than 99% pure
Method/guideline	National Toxicology Program. Toxicology and Carcinogenesis study NTP TR 347
GLP	Yes
Year	1990
Species/strain	B6C3F1 Mice
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	Males: 0, 250 or 500 mg/kg bw/d; Females: 0, 500 or 1000 mg/kg bw/d
Exposure Period	103 weeks
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes
Remarks for Test Conditions	Groups of fifty male and fifty female mice each were administered 0, 250, or 500 mg/kg bw/d or 0, 500 or 1000 mg/kg bw/d <i>d</i> -limonene, respectively, in corn oil by gavage once per day, five days a week for 103 weeks. Animals were housed five per cage and fed ad libitum. The animals were

observed twice per day and weighed once per week for 12 weeks and once per month thereafter. Necropsies were performed on all animals. Histological examinations were performed on all animals dying during the study, all vehicle controls, and all high dose animals. Tissues examined included adrenal glands, brain, cecum, colon, costochondral junction, duodenum, epididymus/seminal vesicles/tunica vaginalis/scrotal sac/prostate/testes or ovaries/uterus, esophagus, eyes, femur or sternbrae or vertebrae including marrow, gallbladder, gross lesions and tissue masses with regional lymph nodes, heart, ileum, jejunum, kidneys, larynx and pharynx, liver, lungs and bronchi, mammary gland, mandibular or mesenteric lymph nodes, nasal cavity and turbinates, oral cavity, pancreas, parathyroids, pituitary gland, preputial or clitoral gland, rectum, salivary glands, sciatic nerve, skin, spinal cord, spleen, stomach, thigh muscle, thymus, thyroid gland, trachea, urinary bladder and Zymbal gland. Tissues examined in low dose groups include liver for female mice.

NOAEL (NOEL)	500 mg/kg bw/d
LOAEL(LOEL)	Undetermined for males; 1000 mg/kg bw/d for female
Toxic Response/effects by Dose Level	<p>Male mice Body weight- No effect. Survival rate-24/50;44/50;39/50. Nonneoplastic effects-No effects. Neoplastic effects-No effects.</p> <p>Female mice Body weight- 10% reduction in high dose group by study end. Survival rate-43/50;44/50;43/50. Nonneoplastic effects-No effects. Neoplastic effects-No effects.</p>
Statistical Evaluation	Yes
Remarks for Results	Mean body weights for female mice administered 1000 mg/kg bw/d <i>d</i> -limonene were generally 5-15% lower than vehicle controls from week 28 to study termination. No treatment related clinical signs were reported for the duration of the study. Survival of the low dose male group was significantly lower than that of the vehicle controls by study termination.
Conclusion Remarks	The authors determined that under the conditions of these 2-year gavage studies there was no evidence of carcinogenic activity of <i>d</i> -limonene for male or female B6C3F1 mice at the dose levels tested.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	National Toxicology Program (NTP) (1990) Carcinogenicity and toxicology studies of <i>d</i> -limonene in F344/N Rats and B6C3F1 mice. NTP-TR-347. U.S. Dept of Health and Human Services. NIH Publication No. 90-2802.

Substance Name	<i>d</i> -Limonene
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CAS No.	5989-27-5
GLP	Ambiguous
Year	1987
Species/strain	F344/N Rats
Sex	Male
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 75, 150, or 300 mg/kg bw/d
Exposure Period	27 days
Frequency of Treatment	Daily (5 days/week)
Control Group	Yes, vehicle only
Remarks for Test Conditions	Groups of five young adult male F344/N rats each were administered <i>d</i> -limonene at dose levels of 0, 75, 150 or 300 mg/kg bw/d five days a week for 27 days. Observations included daily body weight, weekly food intake, liver and kidney weights and light microscopy and histology of liver and kidneys. Rats were examined for hyaline drop formation, granular cast formation and chronic nephrosis. Two-dimensional gel electrophoresis evaluation of protein profiles was conducted on samples of kidneys in the 150 mg/kg dose group killed on day 6.
NOAEL (NOEL)	Less than 75 mg/kg bw/d
LOAEL(LOEL)	75 mg/kg bw/d
Toxic Response/effects by Dose Level	Dose related increases in liver and kidney weights were reported for all dose levels. Renal effects were noted including protein profile changes, hyaline droplet formation, and accumulation of <i>alpha</i> -2-globulin was reported.
Statistical Evaluation	Yes
Conclusion Remarks	Chronic nephrosis was present in all kidneys of treated animals killed on day 27. The authors noted that unlike female rats or higher mammalian species, male rats have anatomical, physiological and biochemical peculiarities involving the proximal convoluted tubule.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Basic data given: comparable to guidelines/standards.
References	Kanerva R.L., Ridder G.M. Lefever F.R. and Alden C.L. (1987) Comparison of short-term renal effects due to oral administration of decalin or <i>d</i> -limonene in young adult male Fischer rats. <i>Fd Chem Toxicol</i> 25, 345-353.
Substance Name	Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck)

CAS No.	8008-57-9
Method/guideline	28-Day Oral Toxicity Study
GLP	Yes
Year	1990
Species/strain	Rat/Sprague Dawley
Sex	Male and Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 240, 600 or 1500 mg/kg bw/d
Exposure Period	30 days
Frequency of Treatment	Daily
Control Group	Yes, vehicle only
Remarks for Test Conditions	Groups of ten rats of each sex were administered 0, 240, 600 or 1500 mg/kg bw/d sweet orange oil in 1% methyl cellulose by gavage daily for 30 days. Observations included survival, clinical observations, body weights, food consumption, clinical pathology, gross pathology, organ weights and histopathology.
NOAEL (NOEL)	Less than 240 mg/kg bw/d
LOAEL(LOEL)	240 mg/kg bw/d
Toxic Response/effects by Dose Level	1500 mg/kg bw/d, Males-
Statistical Evaluation	Yes
Remarks for Results	No treatment related effects were reported for survival, clinical observations, body weights or food consumption. Decreases in glucose related to treatment were reported in the mid-dose females and high dose males and females. Increases in serum albumin and total serum protein were observed in all treated females and the high dose males. Histopathology revealed treatment related lesions in the nonglandular stomach of the high dose males and females and in the kidney of all treated male groups. Kidney weights were also increased in the treated male groups and in the high dose female group. Liver weight increases related to treatment were reported for the high-dose females and all dosed male groups.
Conclusion Remarks	The authors concluded that the NOEL under conditions of this study was less than 240 mg/kg bw/d for both male and female rats. The authors noted that the kidney changes observed in the male rat at all dose levels were expected given the known interaction between limonene and <i>alpha</i> -2-microglobulin. Limonene is the principal constituent (greater than 90%) of orange oil.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.

References

Serota D. (1990) 28-Day oral toxicity study in rats. Unpublished report to FEMA.

Substance Name	Orange peel oil, sweet (Citrus sinensis (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	Immunotoxicity-PFC assay- (Plaque Forming Cell) Assay to sheep red blood cells; Host Resistance Assay- Listeria monocytogenes
GLP	Yes
Year	1994
Species/strain	B6C3F1 mice
Sex	Female
Route of Administration	Oral-Gavage
Doses/concentration Levels	0, 625, 1250 or 2500 mg/kg bw
Exposure Period	5 days
Frequency of Treatment	Daily
Control Group	Yes
Remarks for Test Conditions	Orange oil was administered intragastrically to female B6C3F1 mice daily for 5 days at dose levels of 0, 625, 1250 or 2500 mg/kg bw to determine effects on humoral and cell-mediated immune responses. A host resistance assay (Listeria monocytogenes bacterial challenge) was used to assess cell-mediated immunity while the antibody plaque forming cell response to sheep erythrocytes was used to measure humoral immunity. Other parameters evaluated included body weights, lymphoid organ weights and spleen cellularity but in the absence of modulation of the PFC response, these effects were not considered indicators of immunotoxicity.
NOAEL (NOEL)	2500 mg/kg bw/d
LOAEL(LOEL)	Greater than 2500 mg/kg bw/d
Toxic Response/effects by Dose Level	None
Statistical Evaluation	Yes
Conclusion Remarks	Orange oil had no effects on cell-mediated or humoral immune response at any dose level tested.
Data Qualities Reliabilities	Reliability code 1. Reliable without restriction.
Remarks for Data Reliability	Code 1. Guideline study.
References	Gaworski, C. L., Vollmuth, T. A., Dosier, M. M., Heck, J. D., Dunn, L. T., Ratajczak, H. V. and Thomas, P. T. (1994). An Immunotoxicity Assessment of Food Flavouring Ingredients. Food Chem. Toxicol. 32(5): 409-415.

4.4 Reproductive Toxicity

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene, 90% purity
Method/guideline	Not given
GLP	Ambiguous
Year	1993
Species/Strain	Rat/Wistar
Sex	Female
Route of Administration	Oral
Duration of Test	Approximately 128 days
Doses/Concentration	250, 500, 1000 or 1500 mg/kg bw/d
Premating Exposure period for males	Not applicable
Premating Exposure period for females	27, 28 or 29 days depending upon delivery date
Control Group and Treatment	Yes, vehicle only (corn oil at 2.5 ml/kg bw)
Frequency of Treatment	Daily
Remarks for Test Conditions	<i>beta</i> -Myrcene was administered to female Wistar rats via gavage at dose levels of 0, 250, 500, 1000 or 1500 mg/kg bw/d from the 15th day of gestation until weaning of the offspring which was day 21 postnatal. The vehicle was corn oil. Mortality, weight gain and postnatal development were evaluated. Reproductive capacity was assessed in the exposed offspring upon reaching maturity (120 days).
NOAEL(NOEL)	250 mg/kg bw/d
LOAEL(LOEL)	500 mg/kg bw/d
Appropriate statistical evaluations	One way ANOVA and student t test
Remarks for Results	No adverse effects were noted in the offspring at the lowest dose level tested. Decreased body weight, increased perinatal mortality, and delayed developmental landmarks were noted at the 500, 1000 and 1500 mg/kg bw/d dose levels. Fertility was impaired in female offspring exposed to the two highest doses of <i>beta</i> -myrcene.
Data Reliabilities Qualities	Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Comparable to guideline study with acceptable restrictions.

References Delgado I.F., Nogueira A. C., Souza C.M., Costa M.N., Figueiredo L.H., Mattos A.P., Chahoud I. And Paumgarten F. (1993b) Peri- and postnatal developmental toxicity of *beta*-myrcene in the rat. *Fd Chem Toxic* 31, 623-628.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene, 95% purity
Method/guideline	Not given
GLP	Ambiguous
Year	1998
Species/Strain	Rat/Wistar
Sex	Male and Female
Route of Administration	Oral-Gavage
Duration of Test	Approximately 86 days for female animals; approximately 112 days for male animals
Doses/Concentration	100,300, or 500 mg/kg bw/d
Premating Exposure period for males	91 days
Premating Exposure period for females	21 days
Control Group and Treatment	Yes, vehicle only (peanut oil at 2.5 ml/kg bw)
Frequency of Treatment	Daily
Remarks for Test Conditions	Three experimental groups (15 male and 45 female Wistar rats per group) were administered <i>beta</i> -myrcene dissolved in peanut oil via gavage at dose levels of 0, 100, 300, or 500 mg/kg bw/d. The exposure period was 91 days prior to and during mating the mating period for the males and 21days prior to and during the mating period for females, pregnancy, and lactation until 21 days post parturition. All parent animals were evaluated for weight development, mortality, and toxicity signs. Pregnant females were also evaluated for weight gain, spontaneous abortions, dystocia and prolonged duration of pregnancy. All males were sacrificed and decapitated at the conclusion of mating. One third of the females in each dose group were sacrificed at day 21 of pregnancy. The gravid uterus weight was recorded; resorption and living and dead fetuses were counted. Implantation sites were counted. All fetuses were examined for skeletal abnormalities. After the remaining pregnant females gave birth, the offspring was weighed, and examined for signs of developmental delays, specifically, incisor eruption, fur development, downy hair

NOAEL(NOEL)	development, and eye opening. At weaning on day 21, all mothers were sacrificed and necropsied. 300 mg/kg bw/d
LOAEL(LOEL)	500 mg/kg bw/d
Appropriate statistical evaluations	Yes, one way ANOVA, two tailed student t test
Parental data and F1 as Appropriate	No deaths or signs of toxicity were reported in male rats at any dose level. No statistically significant differences in body weight gain were reported between control and test animals. A slight increase in liver and kidney weights was reported for treated male (highest dose only) and female rats. No morphological alterations of the liver or testis tissue were revealed upon examination. No effects were reported on the number of spermatids in the testis or on the number of spermatozoa in the cauda epididymis. No adverse effects on body weight gain and no other signs of toxicity were observed in treated female rats during the pre-mating or mating periods. No treatment related effects were reported on fertility as measured by the mating index and pregnancy index upon comparison to controls. At the highest dose level, a slight increase in the resorption rate and a parallel decrease in the ratio of live fetuses per implantation site were reported.
Offspring toxicity F1 and F2	Increases in the occurrence of fetal skeleton abnormalities between control and treated groups were reported at the 500 mg/kg bw/d level. No adverse effects were reported on duration of pregnancy, labor, pup mortality, and maternal or offspring weight changes. Slight delays in incisor eruption (300 mg/kg bw/d) and eye opening (100, 300 mg/kg bw/d) were reported but were not dose-related.
Remarks for Results	The authors attributed the increase in skeletal abnormalities at the highest dose level tested to known strain-specific anomalies including increases in dislocated sternums, and lumbar extra ribs.
Conclusion Remarks	The authors concluded that the NOAEL for toxic effects on fertility and general reproductive performance via the oral route was 300 mg <i>beta</i> -myrcene/kg bw/d.
Data Reliabilities Qualities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Paumgarten F.J., De-Carvalho R.R., Souza C.C., Madi K. and Chahoud I. (1998) Study of the effects of <i>beta</i> -Myrcene on rat fertility and general reproductive performance. Braz J Med Biol Res, 31(7).

Substance Name	Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	<i>in vivo</i> Reproductive and Developmental Toxicity Screening Test
GLP	Yes

Year	1989
Species/Strain	Rat/Sprague Dawley
Sex	Female
Route of Administration	Oral-Gavage
Duration of Test	39 days
Doses/Concentration	0, 375, 750 or 1500 mg/kg bw/d
Premating Exposure period for males	Not applicable
Premating Exposure period for females	7 days
Control Group and Treatment	Yes, vehicle only (corn oil)
Frequency of Treatment	Daily
Remarks for Test Conditions	Groups of ten female rats were orally administered sweet orange oil via gavage at dose levels of 0, 375, 750 or 1500 mg/kg bw/d for seven days prior to and through cohabitation, gestation, delivery and a four day lactation period. The vehicle was corn oil. Body weights, food consumption and clinical signs were recorded throughout the observation period. All dams were necropsied and examined for gross lesions on day 25 of presumed gestation for rats not delivering a litter and four days postpartum for rats delivering a litter. Pups delivered were sacrificed on day 4 post partum, any pups dying during the lactation period were necropsied.
NOAEL(NOEL)	Less than 375 mg/kg bw/d
LOAEL(LOEL)	375 mg/kg bw/d
Appropriate statistical evaluations	Yes
Parental data and F1 as Appropriate	No deaths occurred at any dose level. Statistically significant numbers of rats from all three dose groups experienced excess salivation during the pre-mating and gestation periods, and during the lactation period for high-dose animals. The dosed rats had decreased weight gains compared to the control rats during the seven day pre-mating period. Absolute and relative maternal food consumption was significantly decreased for the 750 and 1500 mg/kg bw/d dose groups during the seven day pre-mating period. No treatment related effect on mating performance or fertility was reported at any dose level.
Offspring toxicity F1 and F2	A significant increase in stillbirths and pup deaths was reported for the highest dose group compared to the control group. The treatment with sweet orange oil had no effect on the incidence of malformations or gross lesions in the pups.
Conclusion Remarks	The NOAEL for administration of sweet orange peel oil under the conditions of this study was reported to be less than 375 mg/kg bw/d for maternal toxicity and 750 mg/kg bw/d for offspring development.
Data Reliabilities Qualities	Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Comparable to guideline study with acceptable restrictions.

References Hoberman A.M., Vollmuth T.A., Bennett M.B. and M.S. Christian (1989) An evaluation of food flavoring ingredients using an in vivo reproductive and developmental toxicity screening test. Private communication.

4.5 Developmental/Teratogenicity Toxicity

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Not given
GLP	Ambiguous
Year	1975
Species/strain	Rat/Wistar
Sex	Female
Route of Administration	Oral
Duration of Test	7 days
Doses/concentration Levels	0, 591 or 2869 mg/kg bw/d
Exposure Period	Daily from day 9 to 15 of gestation
Frequency of Treatment	Daily
Control Group and Treatment	Yes
Remarks for Test Conditions	Four groups of twenty Wistar female rats each were administered 0, 591 or 2869 mg/kg bw/d <i>d</i> -limonene on days 9-15 of gestation.
NOAEL(NOEL) maternal toxicity	591 mg/kg bw/d
LOAEL(LOEL) maternal toxicity	2869 mg/kg bw/d
NOAEL (NOEL) developmental toxicity	591 mg/kg bw/d
LOAEL (LOEL) developmental toxicity	2869 mg/kg bw/d
Actual dose received by dose level and sex	Not given
Maternal data with dose level	2869 mg/kg bw/d- maternal body weight decreased, and increased mortality was reported
Fetal Data with Dose Level	2869 mg/kg bw/d-delayed ossification of fetuses metacarpal bone and proximal phalanx; decreased body weights; decreased weights of thymus, spleen, and ovaries.

Appropriate statistical evaluations	Not given
Remarks for Results	At the highest dose level, increases in maternal mortality and decreases in maternal and fetal body weights were reported. Additionally at the highest dose level, delayed ossification of fetal metacarpal bones and proximal phalanx and decreased weights of the thymus, spleen, and ovaries were reported.
Conclusion Remarks	The NOAEL for both maternal and offspring toxicity was reported to be 591 mg/kg bw/d.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Tsuji M., Y.Fujisaki, Y.Arikawa, S.Masuda, S.Kinoshita, A.Okubo, K.Noda, H.Ide and Y.Iwanaga (1975b) Studies on <i>d</i> -limonene as a gallstone solubilizer: Effects on Development of Rat Fetuses and Offsprings. Journal Oyo Yakuri, 10(2), 179.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Remarks for Substance	<i>d</i> -Limonene
Method/guideline	Not given
GLP	Ambiguous
Year	1977
Species/strain	Rabbits/Japanese White
Sex	Female
Route of Administration	Oral
Duration of Test	13 days
Doses/concentration Levels	0, 250, 500 or 1000 mg/kg bw/d
Exposure Period	Daily from day 6 to 18 of gestation
Frequency of Treatment	Daily
Control Group and Treatment	Yes
Remarks for Test Conditions	Pregnant Japanese white rabbits were administered 0, 250, 500 or 1000 mg/kg bw/d <i>d</i> -limonene on days 6 to 18 of gestation.
NOAEL(NOEL) maternal toxicity	250 mg/kg bw/d
LOAEL(LOEL) maternal toxicity	500 mg/kg bw/d
NOAEL (NOEL) developmental toxicity	Greater than 1000 mg/kg bw/d
LOAEL (LOEL) developmental toxicity	Not determined

Actual dose received by dose level and sex	Not given
Maternal data with dose level	1000 mg/kg bw/d-increased mortality; 500 and 1000 mg/kg bw/d-significant decreases in body weight gain and food consumption temporarily observed
Fetal Data with Dose Level	No treatment related effects reported.
Appropriate statistical evaluations	Not given
Remarks for Results	Increased maternal mortality was reported at the highest dose level. Significant decreases in maternal body weight gain and food consumption were temporarily observed at the 500 and 1000 mg/kg bw/d dose levels. No treatment related effects were reported for the offspring.
Conclusion Remarks	The NOAEL for maternal toxicity was 250 mg/kg bw/d. The NOAEL for offspring toxicity was greater than 1000 mg/kg bw/d.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Comparable to guideline study with acceptable restrictions.
References	Kodama R., Okubo A., Sato K., Araki E., Noda K., Ide H., and Ikeda T. (1977a) Studies on <i>d</i> -limonene as a gallstone solubilizer: Effect on development of rabbit fetuses and offsprings. <i>Journal Oyo Yakuri</i> , 13(6), 885-898.

Substance Name	<i>d</i> -Limonene
CAS No.	5989-27-5
Method/guideline	Not given
GLP	Ambiguous
Year	1977
Species/strain	Mice/ICR
Sex	Female
Route of Administration	Oral
Duration of Test	6 days
Doses/concentration Levels	0, 591 or 2363 mg/kg bw/d
Exposure Period	Daily from day 7 to 12 of gestation
Frequency of Treatment	Daily
Control Group and Treatment	Yes
Remarks for Test Conditions	Pregnant ICR mice were administered 0, 591 or 2363 mg/kg bw/d <i>d</i> -limonene on days 7 to 12 of gestation.
NOAEL(NOEL) maternal toxicity	591 mg/kg bw/d
LOAEL(LOEL) maternal toxicity	2363 mg/kg bw/d

NOAEL (NOEL) developmental toxicity	591 mg/kg bw/d
LOAEL (LOEL) developmental toxicity	2363 mg/kg bw/d
Actual dose received by dose level and sex	Not given
Maternal data with dose level	2363 mg/kg bw/d-significant decrease of body weight gain
Fetal Data with Dose Level	2363 mg/kg bw/d-increased incidence of fused ribs compared to control; delayed ossification of some bones, which returned to normal after birth; significant decrease in body weight gain of male offspring
Appropriate statistical evaluations	Not given
Remarks for Results	Significant decreases in body weight gain were reported for pregnant ICR mice administered the highest dose level of <i>d</i> -limonene. In the offspring, increased incidence of fused ribs compared to that of the controls, delayed ossification of some bones and decreased body weight gain were reported at the highest dose level tested.
Conclusion Remarks	The NOAEL for both maternal and offspring toxicity was reported to be 591 mg/kg bw/d.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Comparable to guideline study with acceptable restrictions.
References	Kodama R., Okubo A., Sato K., Araki E., Noda K., Ide H., and Ikeda T. (1977a) Studies on <i>d</i> -limonene as a gallstone solubilizer: Effect on development of mouse fetuses and offsprings. <i>Journal Oyo Yakuri</i> , 13(6), 885-898.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	Not given
GLP	Ambiguous
Year	1993
Species/strain	Rat/Wistar
Sex	Female
Route of Administration	Oral
Duration of Test	10 days
Doses/concentration Levels	0, 250, 500 or 1200 mg/kg bw/d
Exposure Period	Daily from day 6 to 15 of gestation
Frequency of Treatment	Daily

Control Group and Treatment	Yes
Remarks for Test Conditions	Pregnant Wistar rats were administered 0, 250, 500 or 1200 mg/kg bw/d <i>beta</i> -myrcene on gestational days 6-15. The vehicle was corn oil.
NOAEL(NOEL) maternal toxicity	500 mg/kg bw/d
LOAEL(LOEL) maternal toxicity	1200 mg/kg bw/d
NOAEL (NOEL) developmental toxicity	500 mg/kg bw/d
LOAEL (LOEL) developmental toxicity	1200 mg/kg bw/d
Actual dose received by dose level and sex	Not given
Maternal data with dose level	1200 mg/kg bw/d-1/29 maternal deaths. Decreased maternal weight gain.
Fetal Data with Dose Level	1200 mg/kg bw/d-Increased skeletal malformations.
Appropriate statistical evaluations	Yes
Remarks for Results	Decreased maternal weight gain was reported at the 1200 mg/kg bw/d dose. Increased fetal skeletal malformations were reported at the 1200 mg/kg bw/d dose level.
Conclusion Remarks	The NOAEL for both maternal and offspring toxicity was reported to be 500 mg/kg bw/d.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Delgado I.F., Carvalho R.R., Nogueira A., Mattos A.P., Figueiredo L.H., Oliveira S.H.P., Chahoud I., and Paumgarten F.J.R. (1993a) Study on embryo-fetotoxicity of <i>beta</i> -myrcene in the rat. Food Chem Toxic 31, 31-35.

Substance Name	Myrcene
CAS No.	123-35-3
Remarks for Substance	<i>beta</i> -Myrcene
Method/guideline	Not given
GLP	Ambiguous
Year	1993
Species/strain	Rat/Wistar
Sex	Female
Route of Administration	Oral
Duration of Test	Approximately 128 days

Doses/concentration Levels	0, 250, 500, 1000 or 1500 mg/kg bw/d
Exposure Period	27, 28 or 29 days depending upon delivery date
Frequency of Treatment	Daily
Control Group and Treatment	Yes, vehicle only (corn oil at 2.5 ml/kg bw)
Remarks for Test Conditions	<i>beta</i> -Myrcene was administered via gavage to female Wistar rats from pregnancy day 15 until weaning of the offspring which was day 21 postnatal. The vehicle was corn oil. Mortality, weight gain and postnatal development were evaluated. Reproductive capacity was assessed in the exposed offspring upon reaching maturity (120 days).
NOAEL(NOEL) maternal toxicity	500 mg/kg bw/d
LOAEL(LOEL) maternal toxicity	1000 mg/kg bw/d
NOAEL (NOEL) developmental toxicity	250 mg/kg bw/d
LOAEL (LOEL) developmental toxicity	500 mg/kg bw/d
Actual dose received by dose level and sex	Not given
Maternal data with dose level	1500 mg/kg bw/d: 5/15 deaths within first four days of treatment; decreased weight. 1000 mg/kg bw/d: No maternal deaths. Weight deficit observed at term no longer present at parturition.
Fetal Data with Dose Level	500, 1000, or 1500 mg/kg bw/d: decreased body weight, increased perinatal mortality and delayed developmental landmarks; impaired fertility (1000 and 1500 mg/kg bw/d only)
Appropriate statistical evaluations	One way ANOVA and student t test
Remarks for Results	No adverse effects were noted in the offspring at the lowest dose level tested. Decreased body weight, increased perinatal mortality, and delayed developmental landmarks were noted at the 500, 1000 and 1500 mg/kg bw/d dose levels. Fertility was impaired in female offspring exposed to the two highest doses of <i>beta</i> -myrcene.
Conclusion Remarks	The NOAEL for peri- and post natal development was set at 250 mg/kg bw/d.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Comparable to guideline study with acceptable restrictions.
References	Delgado I.F., Nogueira A. C., Souza C.M., Costa M.N., Figueiredo L.H., Mattos A.P., Chahoud I. And Paumgarten F. (1993b) Peri- and postnatal developmental toxicity of <i>beta</i> -myrcene in the rat. <i>Fd Chem Toxic</i> 31, 623-628.

Substance Name	Orange peel oil, sweet (<i>Citrus sinensis</i> (L.) Osbeck)
CAS No.	8008-57-9
Method/guideline	<i>in vivo</i> Reproductive and Developmental Toxicity Screening Test

GLP	Yes
Year	1989
Species/strain	Rat/Sprague Dawley
Sex	Female
Route of Administration	Oral-Gavage
Duration of Test	39 days
Doses/concentration Levels	0, 375, 750 or 1500 mg/kg bw/d
Exposure Period	38 days
Frequency of Treatment	Daily
Control Group and Treatment	Yes, vehicle only (corn oil)
Remarks for Test Conditions	Groups of ten female rats were orally administered sweet orange oil via gavage at dose levels of 0, 375, 750 or 1500, 375 mg/kg bw/d for seven days prior to and through cohabitation, gestation, delivery and a four day lactation period. The vehicle was corn oil. Body weights, food consumption and clinical signs were recorded throughout the observation period. All dams were necropsied and examined for gross lesions on Day 25 of presumed gestation for rats not delivering a litter and four days postpartum for rats delivering a litter. Pups delivered were sacrificed on day 4 post partum, any pups dying during the lactation period were necropsied.
NOAEL(NOEL) maternal toxicity	Less than 375 mg/kg bw/d
LOAEL(LOEL) maternal toxicity	375 mg/kg bw/d
NOAEL (NOEL) developmental toxicity	750 mg/kg bw/d
LOAEL (LOEL) developmental toxicity	1500 mg/kg bw/d
Maternal data with dose level	No deaths occurred at any dose level. Statistically significant numbers of rats from all three dose groups experienced excess salivation during the pre-mating and gestation periods, and during the lactation period for high dose animals. The dosed rats had decreased weight gains compared to the control rats during the seven-day pre-mating period. Absolute and relative maternal food consumption was significantly decreased for the 750 and 1500 mg/kg bw/d dose groups during the seven day pre-mating period. No treatment related effects were reported on maternal body weight, changes in body weight, and absolute and relative feed consumption during the lactation period.
Fetal Data with Dose Level	A significant increase in stillbirths and pup deaths was reported for the highest dose group compared to the control group. The treatment with sweet orange oil had no effect on the incidence of malformations or gross lesions in the pups.
Appropriate statistical evaluations	Yes

Conclusion Remarks	The NOAEL for administration of sweet orange peel oil under the conditions of this study was reported to be less than 375 mg/kg bw/d for maternal toxicity and 750 mg/kg bw/d for offspring development.
Data Qualities Reliabilities	Reliability code 2. Reliable with restriction.
Remarks for Data Reliability	Code 2. Acceptable, well-documented publication/study report, which meets basic scientific principles.
References	Hoberman A.M., Vollmuth T.A., Bennett M.B. and M.S. Christian (1989) An evaluation of food flavoring ingredients using an in vivo reproductive and developmental toxicity screening test. Private communication.